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(54) Title: DOUBLE STRAND COMPOSITIONS COMPRISING DIFFERENTIALLY MODIFIED STRANDS FOR USE IN GENE MODULATION

(57) Abstract: The present invention provides double stranded compositions wherein each strand is modified to have a motif defined by positioning of β -D-ribonucleosides and sugar modified nucleosides. More particularly, the present compositions comprise one strand having an alternating motif and another strand having a hemimer motif, a blockmer motif, a fully modified motif or a positionally modified motif. At least one of the strands has complementarity to a nucleic acid target. The compositions are useful for targeting selected nucleic acid molecules and modulating the expression of one or more genes. In preferred embodiments the compositions of the present invention hybridize to a portion of a target RNA resulting in loss of normal function of the target RNA. The present invention also provides methods for modulating gene expression.



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DOUBLE STRAND COMPOSITIONS COMPRISING DIFFERENTIALLY MODIFIED STRANDS FOR USE IN GENE MODULATION

Cross-Reference to Related Applications

5 The present application: 1) claims benefit to U.S. Provisional Serial No. 60/584,045 filed June 29, 2004, and U.S. Provisional Serial No. 60/607,927 filed September 7, 2004; 2) is a continuation-in-part of U.S. Serial No. 10/859,825 filed June 3, 2004, and U.S. Serial No. 10/946,147 filed September 20, 2004; and 3) is a continuation-in-part of International Serial No. PCT/US2004/017485 filed June 3, 2004, and International Serial No. PCT/US2004/017522 filed
10 June 3, 2004; each of which is incorporated herein by reference in its entirety.

Field of the Invention

 The present invention provides compositions comprising oligomeric compounds that modulate gene expression. In one embodiment, such modulation is via the RNA interference
15 pathway. The modified oligomeric compounds of the invention comprise motifs that can enhance various physical properties and attributes compared to wild type nucleic acids. More particularly, the modification of both strands enables enhancing each strand independently for maximum efficiency for their particular roles in a selected pathway such as the RNAi pathway. The compositions are useful for, for example, targeting selected nucleic acid molecules and
20 modulating the expression of one or more genes. In some embodiments, the compositions of the present invention hybridize to a portion of a target RNA resulting in loss of normal function of the target RNA.

Background of the Invention

25 In many species, introduction of double-stranded RNA (dsRNA) induces potent and specific gene silencing. This phenomenon occurs in both plants and animals and has roles in viral defense and transposon silencing mechanisms. This phenomenon was originally described more than a decade ago by researchers working with the petunia flower. While trying to deepen the purple color of these flowers, Jorgensen et al. introduced a pigment-producing gene under the
30 control of a powerful promoter. Instead of the expected deep purple color, many of the flowers appeared variegated or even white. Jorgensen named the observed phenomenon "cosuppression", since the expression of both the introduced gene and the homologous endogenous gene was suppressed (Napoli et al., Plant Cell, 1990, 2, 279-289; Jorgensen et al., Plant Mol. Biol., 1996, 31, 957-973).

Cosuppression has since been found to occur in many species of plants, fungi, and has been particularly well characterized in *Neurospora crassa*, where it is known as “quelling” (Cogoni et al., *Genes Dev.*, 2000, 10, 638-643; Guru, *Nature*, 2000, 404, 804-808).

The first evidence that dsRNA could lead to gene silencing in animals came from work in the nematode, *C. elegans*. In 1995, researchers Guo and Kemphues were attempting to use antisense RNA to shut down expression of the *par-1* gene in order to assess its function. As expected, injection of the antisense RNA disrupted expression of *par-1*, but quizzically, injection of the sense-strand control also disrupted expression (Guo et al., *Cell*, 1995, 81, 611-620). This result was a puzzle until Fire et al. injected dsRNA (a mixture of both sense and antisense strands) into *C. elegans*. This injection resulted in much more efficient silencing than injection of either the sense or the antisense strands alone. Injection of just a few molecules of dsRNA per cell was sufficient to completely silence the homologous gene’s expression. Furthermore, injection of dsRNA into the gut of the worm caused gene silencing not only throughout the worm, but also in first generation offspring (Fire et al., *Nature*, 1998, 391, 806-811).

The potency of this phenomenon led Timmons and Fire to explore the limits of the dsRNA effects by feeding nematodes bacteria that had been engineered to express dsRNA homologous to the *C. elegans* *unc-22* gene. Surprisingly, these worms developed an *unc-22* null-like phenotype (Timmons et al., *Nature*, 1998, 395, 854; Timmons et al., *Gene*, 2001, 263, 103-112). Further work showed that soaking worms in dsRNA was also able to induce silencing (Tabara et al., *Science*, 1998, 282, 430-431). PCT publication WO 01/48183 discloses methods of inhibiting expression of a target gene in a nematode worm involving feeding to the worm a food organism which is capable of producing a double-stranded RNA structure having a nucleotide sequence substantially identical to a portion of the target gene following ingestion of the food organism by the nematode, or by introducing a DNA capable of producing the double-stranded RNA structure.

The posttranscriptional gene silencing defined in *C. elegans* resulting from exposure to double-stranded RNA (dsRNA) has since been designated as RNA interference (RNAi). This term has come to generalize all forms of gene silencing involving dsRNA leading to the sequence-specific reduction of endogenous targeted mRNA levels; unlike co-suppression, in which transgenic DNA leads to silencing of both the transgene and the endogenous gene.

Introduction of exogenous double-stranded RNA (dsRNA) into *C. elegans* has been shown to specifically and potently disrupt the activity of genes containing homologous sequences. Montgomery et al. suggests that the primary interference effects of dsRNA are post-transcriptional; this conclusion being derived from examination of the primary DNA sequence

after dsRNA-mediated interference a finding of no evidence of alterations followed by studies involving alteration of an upstream operon having no effect on the activity of its downstream gene. These results argue against an effect on initiation or elongation of transcription. Finally they observed by *in situ* hybridization, that dsRNA-mediated interference produced a substantial, although not complete, reduction in accumulation of nascent transcripts in the nucleus, while cytoplasmic accumulation of transcripts was virtually eliminated. These results indicate that the endogenous mRNA is the primary target for interference and suggest a mechanism that degrades the targeted mRNA before translation can occur. It was also found that this mechanism is not dependent on the SMG system, an mRNA surveillance system in *C. elegans* responsible for targeting and destroying aberrant messages. The authors further suggest a model of how dsRNA might function as a catalytic mechanism to target homologous mRNAs for degradation. (Montgomery et al., Proc. Natl. Acad. Sci. U S A, 1998, 95, 15502-15507).

The development of a cell-free system from syncytial blastoderm *Drosophila* embryos that recapitulates many of the features of RNAi has been reported. The interference observed in this reaction is sequence specific, is promoted by dsRNA but not single-stranded RNA, functions by specific mRNA degradation, and requires a minimum length of dsRNA. Furthermore, preincubation of dsRNA potentiates its activity demonstrating that RNAi can be mediated by sequence-specific processes in soluble reactions (Tuschl et al., Genes Dev., 1999, 13, 3191-3197).

In subsequent experiments, Tuschl et al, using the *Drosophila* in vitro system, demonstrated that 21- and 22-nt RNA fragments are the sequence-specific mediators of RNAi. These fragments, which they termed short interfering RNAs (siRNAs) were shown to be generated by an RNase III-like processing reaction from long dsRNA. They also showed that chemically synthesized siRNA duplexes with overhanging 3' ends mediate efficient target RNA cleavage in the *Drosophila* lysate, and that the cleavage site is located near the center of the region spanned by the guiding siRNA. In addition, they suggest that the direction of dsRNA processing determines whether sense or antisense target RNA can be cleaved by the siRNA-protein complex (Elbashir et al., Genes Dev., 2001, 15, 188-200). Further characterization of the suppression of expression of endogenous and heterologous genes caused by the 21-23 nucleotide siRNAs have been investigated in several mammalian cell lines, including human embryonic kidney (293) and HeLa cells (Elbashir et al., Nature, 2001, 411, 494-498).

Tijsterman et al. have shown that, in fact, single-stranded RNA oligomers of antisense polarity can be potent inducers of gene silencing. As is the case for co-suppression, they showed that antisense RNAs act independently of the RNAi genes *rde-1* and *rde-4* but require the

mutator/RNAi gene mut-7 and a putative DEAD box RNA helicase, mut-14. According to the authors, their data favor the hypothesis that gene silencing is accomplished by RNA primer extension using the mRNA as template, leading to dsRNA that is subsequently degraded suggesting that single-stranded RNA oligomers are ultimately responsible for the RNAi phenomenon (Tijsterman et al., Science, 2002, 295, 694-697).

Several other publications have described the structural requirements for the dsRNA trigger required for RNAi activity. Recent reports have indicated that ideal dsRNA sequences are 21nt in length containing 2 nt 3'-end overhangs (Elbashir et al, EMBO (2001), 20, 6877-6887, Sabine Brantl, Biochimica et Biophysica Acta, 2002, 1575, 15-25.) In this system, substitution of the 4 nucleosides from the 3'-end with 2'-deoxynucleosides has been demonstrated to not affect activity. On the other hand, substitution with 2'-deoxynucleosides or 2'-OMe-nucleosides throughout the sequence (sense or antisense) was shown to be deleterious to RNAi activity.

Investigation of the structural requirements for RNA silencing in *C. elegans* has demonstrated modification of the internucleotide linkage (phosphorothioate) to not interfere with activity (Parrish et al., Molecular Cell, 2000, 6, 1077-1087.) It was also shown by Parrish et al., that chemical modification like 2'-amino or 5'-iodouridine are well tolerated in the sense strand but not the antisense strand of the dsRNA suggesting differing roles for the 2 strands in RNAi. Base modification such as guanine to inosine (where one hydrogen bond is lost) has been demonstrated to decrease RNAi activity independently of the position of the modification (sense or antisense). Same "position independent" loss of activity has been observed following the introduction of mismatches in the dsRNA trigger. Some types of modifications, for example introduction of sterically demanding bases such as 5-iodoU, have been shown to be deleterious to RNAi activity when positioned in the antisense strand, whereas modifications positioned in the sense strand were shown to be less detrimental to RNAi activity. As was the case for the 21 nt dsRNA sequences, RNA-DNA heteroduplexes did not serve as triggers for RNAi. However, dsRNA containing 2'-F-2'-deoxynucleosides appeared to be efficient in triggering RNAi response independent of the position (sense or antisense) of the 2'-F-2'-deoxynucleosides.

In one experiment the reduction of gene expression was studied using electroporated dsRNA and a 25mer morpholino in post implantation mouse embryos (Mellitzer et al., Mechanisms of Development, 2002, 118, 57-63). The morpholino oligomer did show activity but was not as effective as the dsRNA.

A number of PCT applications have been published that relate to the RNAi phenomenon. These include: PCT publication WO 00/44895; PCT publication WO 00/49035;

PCT publication WO 00/63364; PCT publication WO 01/36641; PCT publication WO 01/36646; PCT publication WO 99/32619; PCT publication WO 00/44914; PCT publication WO 01/29058; and PCT publication WO 01/75164.

U.S. patents 5,898,031 and 6,107,094 describe certain oligonucleotide having RNA like
5 properties. When hybridized with RNA, these oligonucleotides serve as substrates for a dsRNase enzyme with resultant cleavage of the RNA by the enzyme.

In another published paper (Martinez et al., Cell, 2002, 110, 563-574) it was shown that double stranded as well as single stranded siRNA resides in the RNA-induced silencing complex (RISC) together with eIF2C1 and eIF2C2 (human GERP950 Argonaute proteins. The activity of
10 5'-phosphorylated single stranded siRNA was comparable to the double stranded siRNA in the system studied. In a related study, the inclusion of a 5'-phosphate moiety was shown to enhance activity of siRNA's in vivo in Drosophila embryos (Boutla, et al., Curr. Biol., 2001, 11, 1776-1780). In another study, it was reported that the 5'-phosphate was required for siRNA function in human HeLa cells (Schwarz et al., Molecular Cell, 2002, 10, 537-548).

15 A wide variety of chemical modifications have been made to siRNA compositions to try to enhance properties including stability and potency relative to the unmodified compositions. Much of the early work looked at modification of one strand while keeping the other strand unmodified. More recent work has focused on modification of both strands.

One group is working on modifying both strands of siRNA duplexes such that each
20 strand has an alternating pattern wherein each nucleoside or a block of modified nucleosides is alternating with unmodified β -D-ribonucleosides. The chemical modification used in the modified portion is 2'-OCH₃ modified nucleosides (see European publication EP 1389637 A1, published on February 18, 2004 and PCT publication WO2004015107 published on February 19, 2004).

25 Another group has prepared a number of siRNA constructs with modifications in both strands (see PCT publication WO03/070918 published on August 28, 2003). The constructs disclosed generally have modified nucleosides dispersed in a pattern that is dictated by which strand is being modified and further by the positioning of the purines and pyrimidines in that strand. In general the purines are 2'-OCH₃ or 2'-H and pyrimidines are 2'-F in the antisense
30 strand and the purines are 2'-H and the pyrimidines are 2'-OCH₃ or 2'-F in the sense strand. According to the definitions used in the present application these constructs would appear to be positionally modified as there is no set motif to the substitution pattern and positionally modified can describe a random substitution pattern.

Certain nucleoside compounds having bicyclic sugar moieties are known as locked nucleic acids or LNA (Koshkin et al., Tetrahedron 1998, 54, 3607-3630). These compounds are also referred to in the literature as bicyclic nucleotide analogs (Imanishi *et al.*, International Patent Application WO 98/39352), but this term is also applicable to a genus of compounds that includes other analogs in addition to LNAs. Such modified nucleosides mimic the 3'-endo sugar conformation of native ribonucleosides with the advantage of having enhanced binding affinity and increased resistance to nucleases.

One group recently reported that the incorporation of bicyclic nucleosides, each having a 4'-CH₂-O-2' bridge (LNA) into siRNA duplexes dramatically improved the half life in serum via enhanced nuclease resistance and also increased the duplex stability due to the increased affinity. This effect is seen with a minimum number of LNA's located at specific positions within the siRNA duplex. The placement of LNA's at the 5'-end of the sense strand was shown to reduce the loading of this strand which reduces off target effects (see Elmen et al., Nucleic Acids Res., 2005, 33(1), 439-447).

Some LNAs have a 2'-hydroxyl group linked to the 4' carbon atom of the sugar ring thereby forming a bicyclic sugar moiety. The linkage may be a methylene (-CH₂-)_n group bridging the 2' oxygen atom and the 4' carbon atom wherein n is 1 or 2 (Singh et al., Chem. Commun., 1998, 4, 455-456; Kaneko *et al.*, U.S. Patent Application Publication No.: US 2002/0147332, also see Japanese Patent Application HEI-11-33863, February 12, 1999).

U. S. Patent Application Publication No. 2002/0068708 discloses a number of nucleosides having a variety of bicyclic sugar moieties with the various bridges creating the bicyclic sugar having a variety of configurations and chemical composition.

Braash et al., Biochemistry 2003, 42, 7967-7975 report improved thermal stability of LNA modified siRNA without compromising the efficiency of the siRNA. Grunweller, et. al., Nucleic Acid Research, 2003, 31, 3185-3193 discloses the potency of certain LNA gapmers and siRNAs.

One group has identified a 9 base sequence within an siRNA duplex that elicits a sequence-specific TLR7-dependent immune response in plasmacytoid dendritic cells. The immunostimulation was reduced by incorporating 4 bicyclic nucleosides, each having a 4'-CH₂-O-2' bridge (LNA) at the 3'-end of the sense strand. They also made 5' and both 3' and 5' versions of sense and antisense for incorporation into siRNA duplexes where one strand had the modified nucleosides and the other strand was unmodified (see Hornung et al., 2005, 11(3)I, 263-270).

One group of researchers used expression profiling to perform a genome wide analysis of the efficacy and specificity of siRNA induced silencing of two genes involved in signal transduction (insulin-like growth factor receptor (IGF1R) and mitogen-activated protein kinase 1 (MAPK14 or p38 α). A unique expression profile was produced for each of the 8 siRNAs targeted to MAPK14 and 16 siRNA's targeted to IGF1R indicating that off target effects were highly dependent on the particular sequence. These expression patterns were reproducible for each individual siRNA. The group determined that off target effects were caused by both the antisense strand and the sense strand of siRNA duplexes. There is a need for siRNA's that are designed to preferentially load only the antisense strand thereby reducing the off target effects caused by the sense strand also being loaded into the RISC.

A number of published applications that are commonly assigned with the present application disclose double strand compositions wherein one or both of the strands comprise a particular motif. The motifs include hemimer motifs, blockmer motifs, gapped motifs, fully modified motifs, positionally modified motifs and alternating motifs (see published PCT applications: WO 2004/044133 published May 27, 2004, 3'-endo motifs; WO 2004/113496 published December 29, 2004, 3'-endo motifs; WO 2004/044136 published May 27, 2004, alternating motifs; WO 2004/044140 published May 27, 2004, 2'-modified motifs; WO 2004/043977 published May 27, 2004, 2'-F motifs; WO 2004/043978 published May 27, 2004, 2'-OCH₃ motifs; WO 2004/041889 published May 21, 2004, polycyclic sugar motifs; WO 2004/043979 published May 27, 2004, sugar surrogate motifs; and WO 2004/044138 published May 27, 2004, chimeric motifs; also see published US Application US20050080246 published April 14, 2005).

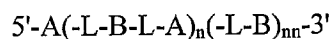
Like the RNase H pathway, the RNA interference pathway of antisense modulation of gene expression is an effective means for modulating the levels of specific gene products and may therefore prove to be uniquely useful in a number of therapeutic, diagnostic, and research applications involving gene silencing. The present invention therefore further provides compositions useful for modulating gene expression pathways, including those relying on an antisense mechanism of action such as RNA interference and dsRNA enzymes as well as non-antisense mechanisms. One having skill in the art, once armed with this disclosure will be able, without undue experimentation, to identify additional compositions for these uses.

Summary of the Invention

In one embodiment, the present invention provides compositions comprising a first oligomeric compound and a second oligomeric compound wherein at least a portion of the first

oligomeric compound is capable of hybridizing with at least a portion of the second oligomeric compound and at least a portion of the first oligomeric compound is complementary to and capable of hybridizing to a selected nucleic acid target. One of the first and second oligomeric compounds comprises nucleosides linked by internucleoside linking groups wherein the linked
 5 nucleosides comprise an alternating motif. The other of the first and second oligomeric compounds comprises nucleosides linked by internucleoside linking groups wherein the linked nucleosides comprise a positionally modified motif, a fully modified motif, a blockmer motif or a hemimer motif. The compositions further comprise one or more optional overhangings, phosphate moieties, conjugate groups or capping groups.

10 The oligomeric compounds comprising an alternating motif include those having the formula:



wherein:

- each L is, independently, an internucleoside linking group;
- 15 each A is a β -D-ribonucleoside or a sugar modified nucleoside;
- each B is a β -D-ribonucleoside or a sugar modified nucleoside;
- n is from about 7 to about 11;
- nn is 0 or 1; and

wherein the sugar groups comprising each A nucleoside are identical, the sugar groups
 20 comprising each B nucleoside are identical, the sugar groups of the A nucleosides are different than the sugar groups of the B nucleosides and at least one of A and B is a sugar modified nucleoside.

In one embodiment, each A or each B is a β -D-ribonucleoside. In another embodiment, each A or each B is a 2'-modified nucleoside wherein the 2'-substituent is selected from halogen,
 25 allyl, amino, azido, O-allyl, O-C₁-C₁₀ alkyl, -OCF₃, O-(CH₂)₂-O-CH₃, 2'-O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n) or O-CH₂-C(=O)-N(R_m)(R_n), where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl. In one embodiment, the 2'-substituent is allyl, O-allyl, O-C₁-C₁₀ alkyl, O-(CH₂)₂-O-CH₃ or 2'-O(CH₂)₂SCH₃ with O-(CH₂)₂-O-CH₃ being particularly suitable.

30 In one embodiment, each A and each B is modified nucleoside. In another embodiment, one of each A and each B comprises 2'-OCH₃ modified nucleosides. In another embodiment, each A and each B comprises 2'-F modified nucleosides.

In one embodiment, the second oligomeric compound comprises an alternating motif and one of each A and each B are β -D-ribonucleosides. In another embodiment, the other of

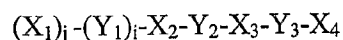
each A and each B comprises 2'-modified nucleosides wherein 2'-substituents include, but are not limited to, allyl, O-allyl, O-C₁-C₁₀ alkyl, O-(CH₂)₂-O-CH₃ or 2'-O(CH₂)₂SCH₃ with O-(CH₂)₂-O-CH₃ being particularly suitable.

In one embodiment, each L is independently a phosphodiester or a phosphorothioate
5 internucleoside linking group.

In one embodiment, one of the first and the second oligomeric compounds comprises a fully modified motif wherein essentially each nucleoside of the oligomeric compound is a sugar modified nucleoside and wherein each sugar modification is the same. In another embodiment, each sugar modified nucleoside is selected from 2'-modified nucleosides, 4'-thio modified
10 nucleosides, 4'-thio-2'-modified nucleosides and nucleosides having bicyclic sugar moieties. In another embodiment, each nucleoside of the fully modified oligomeric compound is a 2'-modified nucleoside wherein 2'-OCH₃ or a 2'-F modified nucleosides are suitable and 2'-OCH₃ modified nucleosides are particularly suitable. In another embodiment, the fully modified oligomeric compound includes one or both of the 3' and 5'-termini having one β-D-ribonucleoside.

In one embodiment, one of the first and second oligomeric compounds comprises a positionally modified wherein the positionally modified motif comprises a continuous sequence of linked nucleosides comprising from about 4 to about 8 regions wherein each region is either a sequence of β-D-ribonucleosides or a sequence of sugar modified nucleosides and wherein the regions are alternating wherein each of the β-D-ribonucleoside regions is flanked on each side by
20 a region of sugar modified nucleosides and each region of sugar modified nucleosides is flanked on each side by a β-D-ribonucleoside region with the exception of regions located the 3' and 5'-termini that will only be flanked on one side and wherein the sugar modified nucleosides are selected from 2'-modified nucleosides, 4'-thio modified nucleosides, 4'-thio-2'-modified nucleosides and nucleosides having bicyclic sugar moieties. In one embodiment, the positionally
25 modified motif comprises from 5 to 7 regions. In another embodiment, the regions of β-D-ribonucleosides comprise from 2 to 8 nucleosides in length. In a further embodiment, the regions of sugar modified nucleosides comprises from 1 to 4 nucleosides in length or from 2 to 3 nucleosides in length.

In one embodiment, oligomeric compounds comprising a positionally modified motif
30 have the formula:



wherein :

X₁ is a sequence of from 1 to about 3 sugar modified nucleosides;

Y₁ is a sequence of from 1 to about 5 β-D-ribonucleosides;

X_2 is a sequence of from 1 to about 3 sugar modified nucleosides;

Y_2 is a sequence of from 2 to about 7 β -D-ribonucleosides;

X_3 is a sequence of from 1 to about 3 sugar modified nucleosides;

Y_3 is a sequence of from 4 to about 6 β -D-ribonucleosides;

5 X_4 is a sequence of from 1 to about 3 sugar modified nucleosides;

i is 0 or 1; and

j is 0 or 1 when i is 1 or 0 when i is 0.

In one embodiment, X_4 is a sequence of 3 sugar modified nucleosides, Y_3 is a sequence of 5 β -D-ribonucleosides, X_3 is a sequence of 2 sugar modified nucleosides; and Y_1 is a sequence of 2 β -D-ribonucleosides. In another embodiment, i is 0 and Y_2 is a sequence of 7 β -D-ribonucleosides. In another embodiment, i is 1, j is 0, Y_2 is a sequence of 2 β -D-ribonucleosides and Y_1 is a sequence of 5 β -D-ribonucleosides. In another embodiment i is 1, j is 1, Y_2 is a sequence of 2 β -D-ribonucleosides, Y_1 is a sequence of 3 β -D-ribonucleosides and X_1 is a sequence of 2 sugar modified nucleosides. In one embodiment each of the sugar modified
15 nucleosides is a 2'-modified nucleoside or a 4'-thio modified nucleoside.

In one embodiment, the first strand of the composition comprises the positional motif. In another embodiment, each internucleoside linking group of the positionally modified oligomeric compound is independently selected from phosphodiester or phosphorothioate.

In one embodiment, each of the first and second oligomeric compounds independently
20 comprises from about 12 to about 30 nucleosides. In a further embodiment, each of the first and second oligomeric compounds independently comprises from about 17 to about 23 nucleosides. In another embodiment, each of the first and second oligomeric compounds independently comprises from about 19 to about 21 nucleosides.

In one embodiment, the first and the second oligomeric compounds form a
25 complementary antisense/sense siRNA duplex.

In one embodiment, the present invention also provides methods of inhibiting gene expression comprising contacting one or more cells, a tissue or an animal with a composition described herein.

In another embodiment, compositions of the invention are used in the preparation of
30 medicaments for inhibiting gene expression in a cell, tissue or animal.

In one embodiment, the present invention also provides a method of inhibiting protein levels in a tumor in an animal comprising contacting the animal with a composition of the invention. In a further embodiment, the contacting is contacting is via intravenous

administration. In even a further embodiment, the tumor is a glioblastoma. In another embodiment, the protein is encoded by the survivin gene.

Description of Embodiments

5 The present invention provides double stranded compositions wherein each strand comprises a motif defined by the location of one or more modified nucleosides or modified and unmodified nucleosides. Motifs derive from the positioning of modified nucleosides relative to other modified or unmodified nucleosides in a strand and are independent of the type of internucleoside linkage, the nucleobase or type of nucleobase e.g. purines or pyrimidines. The
10 compositions of the present invention comprise strands that are differentially modified so that the motifs of each are different. This strategy allows for maximizing the desired properties of each strand independently for their intended role in a process of gene modulation e.g. RNA interference. Tailoring the chemistry and the motif of each strand independently also allows for regionally enhancing each strand. More particularly, the present compositions comprise one
15 strand having an alternating motif and another strand having a hemimer motif, a blockmer motif, a fully modified motif or a positionally modified motif.

 The compositions comprising the various motif combinations of the present invention have been shown to have enhanced properties. The properties that can be enhanced include, but are not limited, to modulation of pharmacokinetic properties through modification of protein
20 binding, protein off-rate, absorption and clearance; modulation of nuclease stability as well as chemical stability; modulation of the binding affinity and specificity of the oligomer (affinity and specificity for enzymes as well as for complementary sequences); and increasing efficacy of RNA cleavage.

 Compositions are provided comprising a first and a second oligomeric compound that
25 are fully or at least partially hybridized to form a duplex region and further comprising a region that is complementary to and hybridizes to a nucleic acid target. It is suitable that such a composition comprise a first oligomeric compound that is an antisense strand having full or partial complementarity to a nucleic acid target and a second oligomeric compound that is a sense strand having one or more regions of complementarity to and forming at least one duplex
30 region with the first oligomeric compound.

 The compositions of the present invention are useful for, for example, modulating gene expression. For example, a targeted cell, group of cells, a tissue or an animal is contacted with a composition of the invention to effect reduction of mRNA that can directly inhibit gene expression. In another embodiment, the reduction of mRNA indirectly upregulates a non-

targeted gene through a pathway that relates the targeted gene to a non-targeted gene. Numerous methods and models for the regulation of genes using compositions of the invention are illustrated in the art and in the example section below.

The compositions of the invention modulate gene expression by hybridizing to a
5 nucleic acid target resulting in loss of its normal function. As used herein, the term "target nucleic acid" or "nucleic acid target" is used for convenience to encompass any nucleic acid capable of being targeted including without limitation DNA, RNA (including pre-mRNA and mRNA or portions thereof) transcribed from such DNA, and also cDNA derived from such RNA. In some embodiments, the target nucleic acid is a messenger RNA. In another
10 embodiment, the degradation of the targeted messenger RNA is facilitated by an activated RISC complex that is formed with compositions of the invention. In another embodiment, the degradation of the targeted messenger RNA is facilitated by a nuclease such as RNaseH.

The present invention provides double stranded compositions wherein one of the strands is useful in, for example, influencing the preferential loading of the opposite strand into
15 the RISC (or cleavage) complex. In particular, the present invention provides oligomeric compounds that comprise chemical modifications in at least one of the strands to drive loading of the opposite strand into the RISC (or cleavage) complex. Such modifications can be used to increase potency of duplex constructs that have been modified to enhance stability. Examples of chemical modifications that drive loading of the second strand are expected to include, but are
20 not limited to, MOE (2'-O(CH₂)₂OCH₃), 2'-O-methyl, -ethyl, -propyl, and -N-methylacetamide. Such modifications can be distributed throughout the strand, or placed at the 5' and/or 3' ends to make a gapmer motif on the sense strand. The compositions are useful for targeting selected nucleic acid molecules and modulating the expression of one or more genes. In some
25 embodiments, the compositions of the present invention hybridize to a portion of a target RNA resulting in loss of normal function of the target RNA.

The present invention provides double stranded compositions wherein one strand comprises an alternating motif and the other strand comprises a hemimer motif, a blockmer motif, a fully modified motif or a positionally modified. Each strand of the compositions of the
30 present invention can be modified to fulfil a particular role in for example the siRNA pathway. Using a different motif in each strand with the same types or different chemical modifications in each strand permits targeting the antisense strand for the RISC complex while inhibiting the incorporation of the sense strand. Within this model each strand can be independently modified such that it is enhanced for its particular role. The antisense strand can be modified at the 5'-end to enhance its role in one region of the RISC while the 3'-end can be modified differentially to

enhance its role in a different region of the RISC. Researchers have been looking at the interaction of the guide sequence and the RISC using various models. Different requirements for the 3'-end, the 5'-end and the region corresponding to the cleavage site of the mRNA are being elucidated through these studies. It has now been shown that the 3'-end of the guide sequence
5 complexes with the PAZ domain while the 5'-end complexes with the Piwi domain (see Song et al., Science, 2004, 305, 1434-1437; Song et al., Nature Structural Biology, 2003, 10(12), 1026-1032; Parker et al., Letters to Nature, 2005, 434, 663-666).

As used in the present invention the term "alternating motif" is meant to include a contiguous sequence of nucleosides comprising two different nucleosides that alternate for
10 essentially the entire sequence of the oligomeric compound. The pattern of alternation can be described by the formula: 5'-A(-L-B-L-A)_n(-L-B)_{nn}-3' where A and B are nucleosides differentiated by having at least different sugar groups, each L is an internucleoside linking group, nn is 0 or 1 and n is from about 7 to about 11. This permits alternating oligomeric compounds from about 17 to about 24 nucleosides in length. This length range is not meant to
15 be limiting as longer and shorter oligomeric compounds are also amenable to the present invention. This formula also allows for even and odd lengths for alternating oligomeric compounds wherein the 3' and 5'-terminal nucleosides are the same (odd) or different (even).

The "A" and "B" nucleosides comprising alternating oligomeric compounds of the present invention are differentiated from each other by having at least different sugar moieties.
20 Each of the A and B nucleosides is selected from β-D-ribonucleosides, 2'-modified nucleosides, 4'-thio modified nucleosides, 4'-thio-2'-modified nucleosides, and bicyclic sugar modified nucleosides. The alternating motif includes the alternation of nucleosides having different sugar groups but is independent from the nucleobase sequence and the internucleoside linkages. The internucleoside linkage can vary at each or selected locations or can be uniform or alternating
25 throughout the oligomeric compound.

Alternating oligomeric compounds of the present invention can be designed to function as the sense or the antisense strand. Alternating 2'-OCH₃/2'-F modified oligomeric compounds have been used as the antisense strand and have shown good activity with a variety of sense strands. One antisense oligomeric compound comprising an alternating motif is a 19mer wherein
30 the A's are 2'-OCH₃ modified nucleosides and the B's are 2'-F modified nucleosides (nn is 0 and n is 9). The resulting alternating oligomeric compound will have a register wherein the 3' and 5'-ends are both 2'-OCH₃ modified nucleosides.

Alternating oligomeric compounds have been designed to function as the sense strand also. The chemistry or register is generally different than for the oligomeric compounds

designed for the antisense strand. When a alternating 2'-F/2'-OCH₃ modified 19mer was paired with the antisense strand in the previous paragraph the preferred orientation was determined to be an offset register wherein both the 3' and 5'-ends of the sense strand were 2'-F modified nucleosides. In a matched register the sugar modifications match between hybridized
5 nucleosides so all the terminal ends of an 19mer would have the same sugar modification. Another alternating motif that has been tested and works in the sense strand is β -D-ribonucleosides alternating with 2'-MOE modified nucleosides.

As used in the present invention the term "fully modified motif" is meant to include a contiguous sequence of sugar modified nucleosides wherein essentially each nucleoside is
10 modified to have the same sugar modification. The compositions of the invention can comprise a fully modified strand as the sense or the antisense strand with the sense strand preferred as the fully modified strand. Suitable sugar modified nucleosides for fully modified strands of the invention include 2'-F, 4'-thio and 2'-OCH₃ with 2'-OCH₃ particularly suitable. In one aspect the 3' and 5'-terminal nucleosides are unmodified.

15 As used in the present invention the term "hemimer motif" is meant to include a sequence of nucleosides that have uniform sugar moieties (identical sugars, modified or unmodified) and wherein one of the 5'-end or the 3'-end has a sequence of from 2 to 12 nucleosides that are sugar modified nucleosides that are different from the other nucleosides in the hemimer modified oligomeric compound. An example of a typical hemimer is a an
20 oligomeric compound comprising β -D-ribonucleosides that have a sequence of sugar modified nucleosides at one of the termini. One hemimer motif includes a sequence of β -D-ribonucleosides having from 2-12 sugar modified nucleosides located at one of the termini. Another hemimer motif includes a sequence of β -D-ribonucleosides having from 2-6 sugar modified nucleosides located at one of the termini with from 2-4 being suitable.

25 As used in the present invention the term "blockmer motif" is meant to include a sequence of nucleosides that have uniform sugars (identical sugars, modified or unmodified) that is internally interrupted by a block of sugar modified nucleosides that are uniformly modified and wherein the modification is different from the other nucleosides. More generally, oligomeric compounds having a blockmer motif comprise a sequence of β -D-ribonucleosides having one
30 internal block of from 2 to 6, or from 2 to 4 sugar modified nucleosides. The internal block region can be at any position within the oligomeric compound as long as it is not at one of the termini which would then make it a hemimer. The base sequence and internucleoside linkages can vary at any position within a blockmer motif.

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As used in the present invention the term "positionally modified motif" is meant to include a sequence of β -D-ribonucleosides wherein the sequence is interrupted by two or more regions comprising from 1 to about 4 sugar modified nucleosides. The positionally modified motif includes internal regions of sugar modified nucleoside and can also include one or both termini. Each particular sugar modification within a region of sugar modified nucleosides is variable with uniform modification desired. The sugar modified regions can have the same sugar modification or can vary such that one region may have a different sugar modification than another region. Positionally modified strands comprise at least two sugar modified regions and at least three when both the 3' and 5'-termini comprise sugar modified regions. Positionally modified oligomeric compounds are distinguished from gapped motifs, hemimer motifs, blockmer motifs and alternating motifs because the pattern of regional substitution defined by any positional motif is not defined by these other motifs. Positionally modified motifs are not determined by the nucleobase sequence or the location or types of internucleoside linkages. The term positionally modified oligomeric compound includes many different specific substitution patterns. A number of these substitution patterns have been prepared and tested in compositions.

Either the antisense or the sense strand of compositions of the present invention can be positionally modified. In one embodiment, the positionally modified strand is designed as the antisense strand. A list of different substitution patterns corresponding to positionally modified oligomeric compounds illustrated in the examples are shown below. This list is meant to be instructive and not limiting.

	ISIS No:Length	Substitution pattern 5'-3'	Modified positions
			underlined are modified from 5'-end
	345838 19mer	5- <u>1</u> -5- <u>1</u> -2- <u>1</u> -2- <u>2</u>	6, 12, 15 and 18-19
	352506 19mer	5- <u>2</u> -2- <u>2</u> -5- <u>3</u>	7-8, 10-11, 17-19
25	352505 19mer	4- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>3</u>	5, 8, 11, 14, 17-19
	xxxxxxx 19mer	4- <u>1</u> -6- <u>1</u> -4- <u>3</u>	5, 12, 17-19
	xxxxxxx 19mer	4- <u>2</u> -4- <u>2</u> -5- <u>2</u>	5-6, 11-12, 18-19
	345839 19mer	4- <u>2</u> -2- <u>2</u> -6- <u>3</u>	5-6, 9-10, 17-19
	xxxxxxx 19mer	3- <u>1</u> -4- <u>1</u> -4- <u>1</u> -3- <u>1</u> -1	4, 9, 14, 18
30	353539 19mer	<u>3</u> -5- <u>1</u> -2- <u>1</u> -4- <u>3</u> *	1-3, 9, 12
	355715 19mer	3- <u>1</u> -4- <u>1</u> -8- <u>1</u> -1	4, 9, 18
	xxxxxxx 19mer	3- <u>1</u> -5- <u>1</u> -7- <u>1</u> -1	4, 10, 18
	384760 19mer	<u>2</u> -7- <u>2</u> -5- <u>3</u> *	1-2, 10-11 and 17-19
	371315 19mer	<u>3</u> -6- <u>2</u> -5- <u>3</u>	1-3, 10-11, 17-19

	353538	19mer	2- <u>1</u> -5- <u>1</u> -2- <u>1</u> -4- <u>3</u>	3, 9, 12, 17-19
	xxxxxx	19mer	2- <u>1</u> -4- <u>1</u> -4- <u>1</u> -4- <u>1</u> -1	3, 8, 13, 18
	336674	20mer	15- <u>1</u> -1- <u>3</u>	16, 18-20
	355712	20mer	4- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>3</u> *	5, 8, 11, 14
5	347348	20mer	<u>3</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>3</u>	1-3, 6, 9, 12, 15, 18-20
	348467	20mer	<u>3</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -2- <u>1</u> -5	1-3, 6, 9, 12, 15
	357278	20mer	3- <u>1</u> -4- <u>1</u> -4- <u>1</u> -3- <u>1</u> -1	4, 9, 14, 18
	xxxxxx	20mer	<u>3</u> -1- <u>1</u> -10- <u>1</u> -1- <u>3</u>	1-3, 5, 16, 18-20
	xxxxxx	20mer	3- <u>1</u> -6- <u>1</u> -7- <u>1</u> -1	4, 11, 19
10	357276	20mer	3- <u>1</u> -3- <u>1</u> -7- <u>1</u> -4	4, 8, 16
	xxxxxx	20mer	3- <u>1</u> -5- <u>2</u> -5- <u>1</u> -3	4, 11, 17
	357275	20mer	3- <u>1</u> -5- <u>1</u> -8- <u>1</u> -1	4, 10, 19
	373424	20mer	<u>3</u> -6- <u>2</u> -5- <u>3</u>	1-3, 11-12, 18-20
	357277	20mer	2- <u>1</u> -5- <u>1</u> -5- <u>1</u> -4- <u>2</u>	3, 9, 15, 20-21
15	345712	20mer	2- <u>2</u> -5- <u>2</u> -5- <u>2</u> -2	3-4, 10-11, 17-18

* indicates that more than one type of sugar modified nucleosides were used in the sugar modified regions.

The term "sugar modified nucleosides" as used in the present invention is intended to include all manner of sugar modifications known in the art. The sugar modified nucleosides can have any heterocyclic base moiety and internucleoside linkage and may include further groups independent from the sugar modification. A group of sugar modified nucleosides includes 2'-modified nucleosides, 4'-thio modified nucleosides, 4'-thio-2'-modified nucleosides, and bicyclic sugar modified nucleosides.

The term "2'-modified nucleoside" as used in the present invention is intended to include all manner of nucleosides having a 2'-substituent group that is other than H and OH. Suitable 2'-substituent groups for 2'-modified nucleosides of the invention include, but are not limited to: halo, allyl, amino, azido, amino, SH, CN, OCN, CF₃, OCF₃, O-, S-, or N(R_m)-alkyl; O-, S-, or N(R_m)-alkenyl; O-, S- or N(R_m)-alkynyl; O-alkylenyl-O-alkyl, alkynyl, alkaryl, aralkyl, O-alkaryl, O-aralkyl, O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n) or O-CH₂-C(=O)-N(R_m)(R_n), where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl. These 2'-substituent groups can be further substituted with substituent groups selected from hydroxyl, amino, alkoxy, carboxy, benzyl, phenyl, nitro (NO₂), thiol, thioalkoxy (S-alkyl), halogen, alkyl, aryl, alkenyl and alkynyl where each R_m is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl.

A list of 2'-substituent groups includes F, -NH₂, N₃, OCF₃, O-CH₃, O(CH₂)₃NH₂, CH₂-CH=CH₂, -O-CH₂-CH=CH₂, OCH₂CH₂OCH₃, 2'-O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n), -O(CH₂)₂O(CH₂)₂N(CH₃)₂, and N-substituted acetamide (O-CH₂-C(=O)-N(R_m)(R_n) where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl. Another list of 2'-substituent groups includes F, OCF₃, O-CH₃, OCH₂CH₂OCH₃, 2'-O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n), -O(CH₂)₂O(CH₂)₂N(CH₃)₂, and N-substituted acetamides (O-CH₂-C(=O)-N(R_m)(R_n) where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl.

Also amenable to the present invention is the manipulation of the stereochemistry of the basic furanose ring system which can be prepared in a number of different configurations. The attachment of the heterocyclic base to the 1'-position can result in the α -anomer (down) or the β -anomer (up). The β -anomer is the anomer found in native DNA and RNA but both forms can be used to prepare oligomeric compounds. A further manipulation can be achieved through the substitution the native form of the furanose with the enantiomeric form e.g. replacement of a native D-furanose with its mirror image enantiomer, the L-furanose. Another way to manipulate the furanose ring system is to prepare stereoisomers such as for example substitution at the 2'-position to give either the ribofuranose (down) or the arabinofuranose (up) or substitution at the 3'-position to give the xylofuranose or by altering the 2', and the 3'-position simultaneously to give a xylofuranose. The use of stereoisomers of the same substituent can give rise to completely different conformational geometry such as for example 2'-F which is 3'-endo in the ribo configuration and 2'-endo in the arabino configuration. The use of different anomeric and stereoisomeric sugars in oligomeric compounds is known in the art and amenable to the present invention.

The term "4'-thio modified nucleoside" is intended to include β -D-ribonucleosides having the 4'-O replaced with 4'-S. The term "4'-thio-2'-modified nucleoside" is intended to include 4'-thio modified nucleosides having the 2'-OH replaced with a 2'-substituent group. The preparation of 4'-thio modified nucleosides is disclosed in publications such as for example U.S. Patent 5,639,837 issued June 17, 1997 and PCT publication WO 2005/027962 published on March 31, 2005. The preparation of 4'-thio-2'-modified nucleosides and their incorporation into oligonucleotides is disclosed in the PCT publication WO 2005/027962 published on March 31, 2005. The 4'-thio-2'-modified nucleosides can be prepared with the same 2'-substituent groups previously mentioned with 2'-OCH₃, 2'-O-(CH₂)₂-OCH₃ and 2'-F are suitable groups.

The term "bicyclic sugar modified nucleoside" is intended to include nucleosides having a second ring formed from the bridging of 2 atoms of the ribose ring. Such bicyclic sugar

modified nucleosides can incorporate a number of different bridging groups that form the second ring and can be formed from different ring carbon atoms on the furanose ring. Bicyclic sugar modified nucleosides wherein the bridge links the 4' and the 2'-carbons and has the formula 4'-(CH₂)_n-O-2' wherein n is 1 or 2 are suitable. The synthesis of bicyclic sugar modified nucleosides is disclosed in US patents 6,268,490, 6,794,499 and published U.S. application 20020147332.

The synthesis and preparation of the bicyclic sugar modified nucleosides wherein the bridge is 4'-CH₂-O-2' having nucleobases selected from adenine, cytosine, guanine, 5-methylcytosine, thymine and uracil, along with their oligomerization, and nucleic acid recognition properties have been described (Koshkin et al., Tetrahedron, 1998, 54, 3607-3630 and WO 98/39352 and WO 99/14226). The L isomer of this bicyclic sugar modified nucleoside has also been prepared (Frieden et al., Nucleic Acids Research, 2003, 31, 6365-6372). The 4'-CH₂-S-2' analog has also been prepared (Kumar et al., Bioorg. Med. Chem. Lett., 1998, 8, 2219-2222), and 2'-amino-LNA (Singh et al., J. Org. Chem., 1998, 63, 10035-10039).

Oligomeric compounds of the present invention can also include one or more terminal phosphate moieties. Terminal phosphate moieties can be located at any terminal nucleoside but are suitable at 5'-terminal nucleosides with the 5'-terminal nucleoside of the antisense strand are also suitable. In one aspect, the terminal phosphate is unmodified having the formula -O-P(=O)(OH)OH. In another aspect, the terminal phosphate is modified such that one or more of the O and OH groups are replaced with H, O, S, N(R) or alkyl where R is H, an amino protecting group or unsubstituted or substituted alkyl.

The term "alkyl," as used herein, refers to a saturated straight or branched hydrocarbon radical containing up to twenty four carbon atoms. Examples of alkyl groups include, but are not limited to, methyl, ethyl, propyl, butyl, isopropyl, n-hexyl, octyl, decyl, dodecyl and the like. Alkyl groups typically include from 1 to about 24 carbon atoms, more typically from 1 to about 12 carbon atoms with from 1 to about 6 carbon atoms are also suitable. Alkyl groups as used herein may optionally include one or more further substituent groups.

The term "alkenyl," as used herein, refers to a straight or branched hydrocarbon chain radical containing up to twenty four carbon atoms having at least one carbon-carbon double bond. Examples of alkenyl groups include, but are not limited to, ethenyl, propenyl, butenyl, 1-methyl-2-buten-1-yl, dienes such as 1,3-butadiene and the like. Alkenyl groups typically include from 2 to about 24 carbon atoms, more typically from 2 to about 12 carbon atoms with from 2 to about 6 carbon atoms are also suitable. Alkenyl groups as used herein may optionally include one or more further substituent groups.

The term "alkynyl," as used herein, refers to a straight or branched hydrocarbon radical containing up to twenty four carbon atoms and having at least one carbon-carbon triple bond. Examples of alkynyl groups include, but are not limited to, ethynyl, 1-propynyl, 1-butylnyl, and the like. Alkynyl groups typically include from 2 to about 24 carbon atoms, more typically from 2 to about 12 carbon atoms with from 2 to about 6 carbon atoms are also suitable. Alkynyl groups as used herein may optionally include one or more further substituent groups.

The term "aliphatic," as used herein, refers to a straight or branched hydrocarbon radical containing up to twenty four carbon atoms wherein the saturation between any two carbon atoms is a single, double or triple bond. An aliphatic group can contain from 1 to about 24 carbon atoms, more typically from 1 to about 12 carbon atoms with from 1 to about 6 carbon atoms being desired. The straight or branched chain of an aliphatic group may be interrupted with one or more heteroatoms that include nitrogen, oxygen, sulfur and phosphorus. Such aliphatic groups interrupted by heteroatoms include without limitation polyalkoxys, such as polyalkylene glycols, polyamines, and polyimines, for example. Aliphatic groups as used herein may optionally include further substituent groups.

The term "alkoxy," as used herein, refers to a radical formed between an alkyl group and an oxygen atom wherein the oxygen atom is used to attach the alkoxy group to a parent molecule. Examples of alkoxy groups include, but are not limited to, methoxy, ethoxy, propoxy, isopropoxy, *n*-butoxy, *sec*-butoxy, *tert*-butoxy, *n*-pentoxy, neopentoxy, *n*-hexoxy and the like. Alkoxy groups as used herein may optionally include further substituent groups.

The terms "halo" and "halogen," as used herein, refer to an atom selected from fluorine, chlorine, bromine and iodine.

The terms "aryl" and "aromatic," as used herein, refer to a mono- or polycyclic carbocyclic ring system radical having one or more aromatic rings. Examples of aryl groups include, but not limited to, phenyl, naphthyl, tetrahydronaphthyl, indanyl, idenyl and the like. Aryl groups as used herein may optionally include further substituent groups.

The term "heterocyclic," as used herein, refers to a radical mono-, or poly-cyclic ring system that includes at least one heteroatom and is unsaturated, partially saturated or fully saturated, thereby including heteroaryl groups. Heterocyclic is also meant to include fused ring systems wherein one or more of the fused rings contain no heteroatoms. A heterocyclic group typically includes at least one atom selected from sulfur, nitrogen or oxygen. Examples of heterocyclic groups include, [1,3]dioxolane, pyrrolidinyl, pyrazolinyl, pyrazolidinyl, imidazolinyl, imidazolidinyl, piperidinyl, piperazinyl, oxazolidinyl, isoxazolidinyl, morpholinyl,

thiazolidinyl, isothiazolidinyl, quinoxalinyl, pyridazinonyl, tetrahydrofuryl and the like.

Heterocyclic groups as used herein may optionally include further substituent groups.

The terms "substituent and substituent group," as used herein, are meant to include groups that are typically added to other groups or parent compounds to enhance desired
5 properties or give desired effects. Substituent groups can be protected or unprotected and can be added to one available site or to many available sites in a parent compound. Substituent groups may also be further substituted with other substituent groups and may be attached directly or via a linking group such as an alkyl or hydrocarbyl group to the parent compound. Such substituent groups include without limitation, halogen, hydroxyl, alkyl, alkenyl, alkynyl, acyl ($-C(O)R_a$),
10 carboxyl ($-C(O)O-R_a$), aliphatic, alicyclic, alkoxy, substituted oxo ($-O-R_a$), aryl, aralkyl, heterocyclic, heteroaryl, heteroarylalkyl, amino ($-NR_bR_c$), imino ($=NR_b$), amido ($-C(O)NR_bR_c$ or $-N(R_b)C(O)R_a$), azido ($-N_3$), nitro ($-NO_2$), cyano ($-CN$), carbamido ($-OC(O)NR_bR_c$ or $-N(R_b)C(O)OR_a$), ureido ($-N(R_b)C(O)NR_bR_c$), thioureido ($-N(R_b)C(S)NR_bR_c$), guanidinyl ($-N(R_b)C(=NR_b)NR_bR_c$), amidinyl ($-C(=NR_b)NR_bR_c$
15 or $-N(R_b)C(NR_b)R_a$), thiol ($-SR_b$), sulfinyl ($-S(O)R_b$), sulfonyl ($-S(O)_2R_b$) and sulfonamidyl ($-S(O)_2NR_bR_c$ or $-N(R_b)S(O)_2R_b$). Wherein each R_a , R_b and R_c is a further substituent group which can be without limitation alkyl, alkenyl, alkynyl, aliphatic, alkoxy, acyl, aryl, aralkyl, heteroaryl, alicyclic, heterocyclic and heteroarylalkyl.

The term "protecting group," as used herein, refers to a labile chemical moiety which is
20 known in the art to protect reactive groups including without limitation, hydroxyl, amino and thiol groups, against undesired reactions during synthetic procedures. Protecting groups are typically used selectively and/or orthogonally to protect sites during reactions at other reactive sites and can then be removed to leave the unprotected group as is or available for further reactions. Protecting groups as known in the art are described generally in Greene and Wuts,
25 Protective Groups in Organic Synthesis, 3rd edition, John Wiley & Sons, New York (1999).

Examples of hydroxyl protecting groups include, but are not limited to, benzyloxy-carbonyl, 4-nitrobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-methoxybenzyloxycarbonyl, methoxycarbonyl, tert-butoxycarbonyl (BOC), isopropoxycarbonyl, diphenylmethoxycarbonyl, 2,2,2-trichloroethoxycarbonyl, 2-(trimethylsilyl)ethoxycarbonyl, 2-furfuryloxycarbonyl,
30 allyloxycarbonyl (Alloc), acetyl (Ac), formyl, chloroacetyl, trifluoroacetyl, methoxyacetyl, phenoxyacetyl, benzoyl (Bz), methyl, t-butyl, 2,2,2-trichloroethyl, 2-trimethylsilyl ethyl, 1,1-dimethyl-2-propenyl, 3-methyl-3-butenyl, allyl, benzyl (Bn), para-methoxybenzyl, diphenylmethyl, triphenylmethyl (trityl), 4,4'-dimethoxytriphenylmethyl (DMT), substituted or unsubstituted 9-(9-phenyl)xanthenyl (pixyl), tetrahydrofuryl, methoxymethyl,

methylthiomethyl, benzyloxymethyl, 2,2,2-trichloroethoxymethyl, 2-(trimethylsilyl)ethoxymethyl, methanesulfonyl, para-toluenesulfonyl, trimethylsilyl, triethylsilyl, triisopropylsilyl, and the like. Suitable hydroxyl protecting groups for the present invention are DMT and substituted or unsubstituted pixyl.

5 Examples of amino protecting groups include, but are not limited to, *t*-butoxycarbonyl (BOC), 9-fluorenylmethoxycarbonyl (Fmoc), benzyloxycarbonyl, and the like. Examples of thiol protecting groups include, but are not limited to, triphenylmethyl (Trt), benzyl (Bn), and the like.

 The synthesized oligomeric compounds can be separated from a reaction mixture and
10 further purified by a method such as column chromatography, high pressure liquid chromatography, precipitation, or recrystallization. Further methods of synthesizing the compounds of the formulae herein will be evident to those of ordinary skill in the art. Additionally, the various synthetic steps may be performed in an alternate sequence or order to give the desired compounds. Synthetic chemistry transformations and protecting group
15 methodologies (protection and deprotection) useful in synthesizing the compounds described herein are known in the art and include, for example, those such as described in R. Larock, Comprehensive Organic Transformations, VCH Publishers (1989); T. W. Greene and P. G. M. Wuts, Protective Groups in Organic Synthesis, 2d. Ed., John Wiley and Sons (1991); L. Fieser and M. Fieser, Fieser and Fieser's Reagents for Organic Synthesis, John Wiley and Sons (1994);
20 and L. Paquette, ed., Encyclopedia of Reagents for Organic Synthesis, John Wiley and Sons (1995), and subsequent editions thereof.

 The compounds described herein contain one or more asymmetric centers and thus give rise to enantiomers, diastereomers, and other stereoisomeric forms that may be defined, in terms of absolute stereochemistry, as (R)- or (S)-, or as (D)- or (L)- for amino acids. The present
25 invention is meant to include all such possible isomers, as well as their racemic and optically pure forms. Optical isomers may be prepared from their respective optically active precursors by the procedures described above, or by resolving the racemic mixtures. The resolution can be carried out in the presence of a resolving agent, by chromatography or by repeated crystallization or by some combination of these techniques which are known to those skilled in the art. Further
30 details regarding resolutions can be found in Jacques, et al., Enantiomers, Racemates, and Resolutions (John Wiley & Sons, 1981). When the compounds described herein contain olefinic double bonds, other unsaturation, or other centers of geometric asymmetry, and unless specified otherwise, it is intended that the compounds include both E and Z geometric isomers or cis- and trans-isomers. Likewise, all tautomeric forms are also intended to be included. The

configuration of any carbon-carbon double bond appearing herein is selected for convenience only and is not intended to designate a particular configuration unless the text so states; thus a carbon-carbon double bond or carbon-heteroatom double bond depicted arbitrarily herein as trans may be cis, trans, or a mixture of the two in any proportion.

5 The term “nucleoside,” as used herein, refers to a base-sugar combination. The base portion of the nucleoside is normally a heterocyclic base moiety. The two most common classes of such heterocyclic bases are purines and pyrimidines. Nucleotides are nucleosides that further include a phosphate group covalently linked to the sugar portion of the nucleoside. For those nucleosides that include a pentofuranosyl sugar, the phosphate group can be linked to either the
10 2', 3' or 5' hydroxyl moiety of the sugar. The term nucleoside is intended to include both modified and unmodified nucleosides. Within the oligonucleotide structure, the phosphate groups are commonly referred to as forming the backbone of the oligomeric compound. In forming oligonucleotides, the phosphate groups covalently link adjacent nucleosides to one another to form a linear polymeric compound. The normal internucleoside linkage of RNA and
15 DNA is a 3' to 5' phosphodiester linkage.

In the context of this invention, the term “oligonucleoside” refers to a sequence of nucleosides that are joined by internucleoside linkages that do not have phosphorus atoms. Internucleoside linkages of this type are further described in the “modified internucleoside linkage” section below.

20 The term “oligonucleotide,” as used herein, refers to an oligomer or polymer of ribonucleic acid (RNA) or deoxyribonucleic acid (DNA) composed of naturally occurring nucleobases, sugars and phosphodiester internucleoside linkages.

The terms “oligomer” and “oligomeric compound,” as used herein, refer to a plurality of naturally occurring and/or non-naturally occurring nucleosides, joined together with
25 internucleoside linking groups in a specific sequence. At least some of the oligomeric compounds can be capable of hybridizing a region of a target nucleic acid. Included in the terms “oligomer” and “oligomeric compound” are oligonucleotides, oligonucleotide analogs, oligonucleotide mimetics, oligonucleosides and chimeric combinations of these. As such the term oligomeric compound is broader than the term “oligonucleotide,” including all oligomers
30 having all manner of modifications including but not limited to those known in the art. Oligomeric compounds are typically structurally distinguishable from, yet functionally interchangeable with, naturally-occurring or synthetic wild-type oligonucleotides. Thus, oligomeric compounds include all such structures that function effectively to mimic the structure and/or function of a desired RNA or DNA strand, for example, by hybridizing to a target. Such

non-naturally occurring oligonucleotides are often desired over the naturally occurring forms because they often have enhanced properties, such as for example, enhanced cellular uptake, enhanced affinity for nucleic acid target and increased stability in the presence of nucleases.

Oligomeric compounds can include compositions comprising double-stranded constructs such as, for example, two oligomeric compounds forming a double stranded hybridized construct or a single strand with sufficient self complementarity to allow for hybridization and formation of a fully or partially double-stranded compound. In one embodiment of the invention, double-stranded oligomeric compounds encompass short interfering RNAs (siRNAs). As used herein, the term "siRNA" is defined as a double-stranded construct comprising a first and second strand and having a central complementary portion between the first and second strands and terminal portions that are optionally complementary between the first and second strands or with a target nucleic acid. Each strand in the complex may have a length or from about 12 to about 24 nucleosides and may further comprise a central complementary portion having one of these defined lengths. Each strand may further comprise a terminal unhybridized portion having from 1 to about 6 nucleobases in length. The siRNAs may also have no terminal portions (overhangs) which is referred to as being blunt ended. The two strands of an siRNA can be linked internally leaving free 3' or 5' termini or can be linked to form a continuous hairpin structure or loop. The hairpin structure may contain an overhang on either the 5' or 3' terminus producing an extension of single-stranded character.

In one embodiment of the invention, compositions comprising double-stranded constructs are canonical siRNAs. As used herein, the term "canonical siRNA" is defined as a double-stranded oligomeric compound having a first strand and a second strand each strand being 21 nucleobases in length with the strands being complementary over 19 nucleobases and having on each 3' termini of each strand a deoxy thymidine dimer (dTdT) which in the double-stranded compound acts as a 3' overhang. In another aspect compositions comprise double-stranded constructs having overhangs may be of varying lengths with overhangs of varying lengths and may include compositions wherein only one strand has an overhang.

In another embodiment, compositions comprising double-stranded constructs are blunt-ended siRNAs. As used herein the term "blunt-ended siRNA" is defined as an siRNA having no terminal overhangs. That is, at least one end of the double-stranded constructs is blunt. siRNAs that have one or more overhangs or that are blunt act to elicit dsRNAse enzymes and trigger the recruitment or activation of the RNAi antisense mechanism. In a further embodiment, single-stranded RNAi (ssRNAi) compounds that act via the RNAi antisense mechanism are contemplated.

Further modifications can be made to the double-stranded compounds and may include conjugate groups attached to one or more of the termini, selected nucleobase positions, sugar positions or to one of the internucleoside linkages. Alternatively, the two strands can be linked via a non-nucleic acid moiety or linker group. When formed from only one strand, dsRNA can
5 take the form of a self-complementary hairpin-type molecule that doubles back on itself to form a duplex. Thus, the dsRNAs can be fully or partially double-stranded. When formed from two strands, or a single strand that takes the form of a self-complementary hairpin-type molecule doubled back on itself to form a duplex, the two strands (or duplex-forming regions of a single strand) are complementary RNA strands that base pair in Watson-Crick fashion.

10 The oligomeric compounds in accordance with this invention comprise from about 8 to about 80 nucleobases (i.e. from about 8 to about 80 linked nucleosides/monomeric subunits, or up to 80 linked nucleosides/monomeric subunits). One of ordinary skill in the art will appreciate that the invention embodies oligomeric compounds of 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45,
15 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, or 80 nucleobases in length, or any range therewithin.

In one embodiment, the oligomeric compounds of the invention are 10 to 50 nucleobases in length, or up to 50 nucleobases in length. One having ordinary skill in the art will appreciate that this embodies oligomeric compounds of 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,
20 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, or 50 nucleobases in length, or any range therewithin.

In another embodiment, the oligomeric compounds of the invention are 12 to 30 nucleobases in length, or up to 30 nucleobases in length. One having ordinary skill in the art will appreciate that this embodies oligomeric compounds of 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22,
25 23, 24, 25, 26, 27, 28, 29, or 30 nucleobases in length, or any range therewithin.

In another embodiment, the oligomeric compounds of the invention are 17 to 23 nucleobases in length, or up to 23 nucleobases in length. One having ordinary skill in the art will appreciate that this embodies oligomeric compounds of 17, 18, 19, 20, 21, 22 or 23 nucleobases in length, or any range therewithin.

30 In another embodiment, the oligomeric compounds of the invention are 19 to 21 nucleobases in length, or up to 21 nucleobases in length. One having ordinary skill in the art will appreciate that this embodies oligomeric compounds of 19, 20 or 21 nucleobases in length, or any range therewithin.

As used herein the term "heterocyclic base moiety" refers to nucleobases and modified or substitute nucleobases used to form nucleosides of the invention. The term "heterocyclic base moiety" includes unmodified nucleobases such as the native purine bases adenine (A) and guanine (G), and the pyrimidine bases thymine (T), cytosine (C) and uracil (U). The term is also intended to include all manner of modified or substitute nucleobases including but not limited to synthetic and natural nucleobases such as xanthine, hypoxanthine, 2-aminopyridine and 2-pyridone, 5-methylcytosine (5-me-C), 5-hydroxymethylenyl cytosine, 2-amino and 2-fluoro-adenine, 2-propyl and other alkyl derivatives of adenine and guanine, 2-thio cytosine, uracil, thymine, 3-deaza guanine and adenine, 4-thiouracil, 5-uracil (pseudouracil), 5-propynyl (-C≡C-CH₃) uracil and cytosine and other alkynyl derivatives of pyrimidine bases, 5-halo particularly 5-bromo, 5-trifluoromethyl and other 5-substituted uracils and cytosines, 6-methyl and other alkyl derivatives of adenine and guanine, 6-azo uracil, cytosine and thymine, 7-methyl adenine and guanine, 7-deaza adenine and guanine, 8-halo, 8-amino, 8-aza, 8-thio, 8-thioalkyl, 8-hydroxyl and other 8-substituted adenines and guanines, universal bases, hydrophobic bases, promiscuous bases, size-expanded bases, and fluorinated bases as defined herein. Further modified nucleobases include tricyclic pyrimidines such as phenoxazine cytidine (1H-pyrimido[5,4-b][1,4]benzoxazin-2(3H)-one) and phenothiazine cytidine (1H-pyrimido[5,4-b][1,4]benzothiazin-2(3H)-one).

Further nucleobases (and nucleosides comprising the nucleobases) include those disclosed in US Patent No. 3,687,808, those disclosed in *The Concise Encyclopedia Of Polymer Science And Engineering*, pages 858-859, Kroschwitz, J.I., ed. John Wiley & Sons, 1990, those disclosed by Englisch *et al.*, *Angewandte Chemie*, International Edition, 1991, 30, 613, those disclosed in Limbach *et al.*, *Nucleic Acids Research*, 1994, 22(12), 2183-2196, and those disclosed by Sanghvi, Y.S., Chapter 15, *Antisense Research and Applications*, pages 289-302, Crooke, S.T. and Lebleu, B., ed., CRC Press, 1993.

Certain of these nucleobases are particularly useful for increasing the binding affinity of the oligomeric compounds of the invention. These include 5-substituted pyrimidines, 6-azapyrimidines and N-2, N-6 and O-6 substituted purines, including 2-aminopropyl-adenine, 5-propynyluracil and 5-propynylcytosine. 5-methylcytosine substitutions have been shown to increase nucleic acid duplex stability by 0.6-1.2°C (Sanghvi, Y.S., Crooke, S.T. and Lebleu, B., eds., *Antisense Research and Applications*, CRC Press, Boca Raton, 1993, pp. 276-278) and are especially useful when combined with 2'-O-methoxyethyl (2'-MOE) sugar modifications.

Representative U.S. patents that teach the preparation of certain of the above noted modified nucleobases as well as other modified nucleobases include, but are not limited to, the

above noted U.S. 3,687,808, as well as U.S.: 4,845,205; 5,130,302; 5,134,066; 5,175,273; 5,367,066; 5,432,272; 5,457,187; 5,459,255; 5,484,908; 5,502,177; 5,525,711; 5,552,540; 5,587,469; 5,594,121; 5,596,091; 5,614,617; 5,645,985; 5,830,653; 5,763,588; 6,005,096; 5,681,941, and 5,750,692.

- 5 The term "universal base" as used herein, refers to a moiety that may be substituted for any base. The universal base need not contribute to hybridization, but should not significantly detract from hybridization and typically refers to a monomer in a first sequence that can pair with a naturally occurring base, i.e A, C, G, T or U at a corresponding position in a second sequence of a duplex in which one or more of the following is true: (1) there is essentially no pairing
- 10 (hybridization) between the two; or (2) the pairing between them occurs non-discriminant with the universal base hybridizing one or more of the the naturally occurring bases and without significant destabilization of the duplex. Exemplary universal bases include, without limitation, inosine, 5-nitroindole and 4-nitrobenzimidazole. For further examples and descriptions of universal bases see Survey and summary: the applications of universal DNA base analogs.
- 15 Loakes, Nucleic Acids Research, 2001, 29, 12, 2437-2447.

- The term "promiscuous base" as used herein, refers to a monomer in a first sequence that can pair with a naturally occurring base, i.e A, C, G, T or U at a corresponding position in a second sequence of a duplex in which the promiscuous base can pair non-discriminantly with more than one of the naturally occurring bases, i.e. A, C, G, T, U. Non-limiting examples of
- 20 promiscuous bases are 6H,8H-3,4-dihydropyrimido[4,5-c][1,2]oxazin-7-one and N⁶-methoxy-2,6-diaminopurine, shown below. For further information, see Polymerase recognition of synthetic oligodeoxyribonucleotides incorporating degenerate pyrimidine and purine bases. Hill, et al., Proc. Natl. Acad. Sci., 1998, 95, 4258-4263.

- Examples of G-clamps include substituted phenoxazine cytidine (e.g. 9-(2-aminoethoxy)-H-pyrimido[5,4-b][1,4]benzoxazin-2(3H)-one), carbazole cytidine (2H-pyrimido[4,5-b]indol-2-one) and pyridoindole cytidine (H-pyrido[3',2':4,5]pyrrolo[2,3-d]pyrimidin-2-one).
- 25

- Representative cytosine analogs that make 3 hydrogen bonds with a guanosine in a second oligonucleotide include 1,3-diazaphenoxazine-2-one (Kurchavov et al., Nucleosides and
- 30 Nucleotides, 1997, 16, 1837-1846), 1,3-diazaphenothiazine-2-one (Lin et al., J. Am. Chem. Soc. 1995, 117, 3873-3874) and 6,7,8,9-tetrafluoro-1,3-diazaphenoxazine-2-one (Wang et al., Tetrahedron Lett. 1998, 39, 8385-8388). When incorporated into oligonucleotides these base modifications hybridized with complementary guanine (the latter also hybridized with adenine)

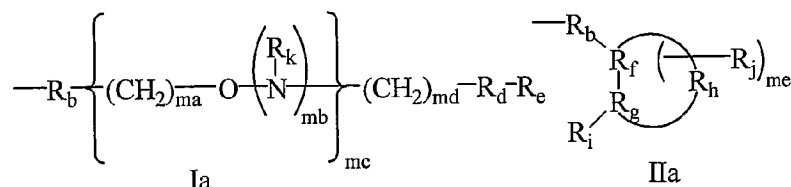
and enhanced helical thermal stability by extended stacking interactions (see U.S. Serial Number 10/013,295).

Oligomeric compounds of the invention may also contain one or more substituted sugar moieties such as the 2'-modified sugars discussed. A more comprehensive but not limiting list of sugar substituent groups includes: OH; F; O-, S-, or N-alkyl; O-, S-, or N-alkenyl; O-, S- or N-alkynyl; or O-alkyl-O-alkyl, wherein the alkyl, alkenyl and alkynyl may be substituted or unsubstituted C₁ to C₁₀ alkyl or C₂ to C₁₀ alkenyl and alkynyl. Particularly suitable are O((CH₂)_nO)_mCH₃, O(CH₂)_nOCH₃, O(CH₂)_nNH₂, O(CH₂)_nCH₃, O(CH₂)_nONH₂, and O(CH₂)_nON((CH₂)_nCH₃)₂, where n and m are from 1 to about 10. Some oligonucleotides comprise a sugar substituent group selected from: C₁ to C₁₀ lower alkyl, substituted lower alkyl, alkenyl, alkynyl, alkaryl, aralkyl, O-alkaryl or O-aralkyl, SH, SCH₃, OCN, Cl, Br, CN, CF₃, OCF₃, SOCH₃, SO₂CH₃, ONO₂, NO₂, N₃, NH₂, heterocycloalkyl, heterocycloalkaryl, aminoalkylamino, polyalkylamino, substituted silyl, an RNA cleaving group, a reporter group, an intercalator, a group for improving the pharmacokinetic properties of an oligonucleotide, or a group for improving the pharmacodynamic properties of an oligonucleotide, and other substituents having similar properties.

One modification includes 2'-methoxyethoxy (2'-O-CH₂CH₂OCH₃, also known as 2'-O-(2-methoxyethyl) or 2'-MOE) (Martin et al., *Helv. Chim. Acta*, 1995, 78, 486-504) i.e., an alkoxyalkoxy group. One modification includes 2'-dimethylaminoethoxyethoxy, i.e., a O(CH₂)₂ON(CH₃)₂ group, also known as 2'-DMAOE, as described in examples hereinbelow, and 2'-dimethylaminoethoxyethoxy (also known in the art as 2'-O-dimethyl-amino-ethoxy-ethyl or 2'-DMAEOE), i.e., 2'-O-CH₂-O-CH₂-N(CH₃)₂.

Other sugar substituent groups include methoxy (-O-CH₃), aminopropoxy (-OCH₂CH₂CH₂NH₂), allyl (-CH₂-CH=CH₂), -O-allyl (-O-CH₂-CH=CH₂) and fluoro (F). 2'-Sugar substituent groups may be in the arabino (up) position or ribo (down) position. One 2'-arabino modification is 2'-F. Similar modifications may also be made at other positions on the oligomeric compound, particularly the 3' position of the sugar on the 3' terminal nucleoside or in 2'-5' linked oligonucleotides and the 5' position of 5' terminal nucleotide. Oligomeric compounds may also have sugar mimetics such as cyclobutyl moieties in place of the pentofuranosyl sugar. Representative U.S. patents that teach the preparation of such modified sugar structures include, but are not limited to, U.S.: 4,981,957; 5,118,800; 5,319,080; 5,359,044; 5,393,878; 5,446,137; 5,466,786; 5,514,785; 5,519,134; 5,567,811; 5,576,427; 5,591,722; 5,597,909; 5,610,300; 5,627,053; 5,639,873; 5,646,265; 5,658,873; 5,670,633; 5,792,747; and 5,700,920.

Representative sugar substituent groups include groups of formula Ia or IIa:



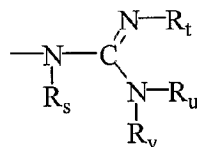
wherein:

R_b is O, S or NH;

5 R_d is a single bond, O, S or C(=O);

R_e is C₁-C₁₀ alkyl, N(R_k)(R_m), N(R_k)(R_n), N=C(R_p)(R_q), N=C(R_p)(R_r) or has formula

III_a;



IIIa

R_p and R_q are each independently hydrogen or C₁-C₁₀ alkyl;

10 R_r is -R_x-R_y;

each R_s, R_t, R_u and R_v is, independently, hydrogen, C(O)R_w, substituted or unsubstituted C₁-C₁₀ alkyl, substituted or unsubstituted C₂-C₁₀ alkenyl, substituted or unsubstituted C₂-C₁₀ alkynyl, alkylsulfonyl, arylsulfonyl, a chemical functional group or a conjugate group, wherein the substituent groups are selected from hydroxyl, amino, alkoxy,

15 carboxy, benzyl, phenyl, nitro, thiol, thioalkoxy, halogen, alkyl, aryl, alkenyl and alkynyl;

or optionally, R_u and R_v, together form a phthalimido moiety with the nitrogen atom to which they are attached;

each R_w is, independently, substituted or unsubstituted C₁-C₁₀ alkyl, trifluoromethyl, cyanoethoxy, methoxy, ethoxy, t-butoxy, allyloxy, 9-fluorenylmethoxy, 2-(trimethylsilyl)-

20 ethoxy, 2,2,2-trichloroethoxy, benzyloxy, butyryl, iso-butyryl, phenyl or aryl;

R_k is hydrogen, a nitrogen protecting group or -R_x-R_y;

R_p is hydrogen, a nitrogen protecting group or -R_x-R_y;

R_x is a bond or a linking moiety;

R_y is a chemical functional group, a conjugate group or a solid support medium;

25 each R_m and R_n is, independently, H, a nitrogen protecting group, substituted or

unsubstituted C₁-C₁₀ alkyl, substituted or unsubstituted C₂-C₁₀ alkenyl, substituted or unsubstituted C₂-C₁₀ alkynyl, wherein the substituent groups are selected from hydroxyl, amino,

alkoxy, carboxy, benzyl, phenyl, nitro, thiol, thioalkoxy, halogen, alkyl, aryl, alkenyl, alkynyl; NH_3^+ , $\text{N}(\text{R}_u)(\text{R}_v)$, guanidino and acyl where the acyl is an acid amide or an ester;

or R_m and R_n , together, are a nitrogen protecting group, are joined in a ring structure that optionally includes an additional heteroatom selected from N and O or are a chemical

5 functional group;

R_i is OR_z , SR_z , or $\text{N}(\text{R}_z)_2$;

each R_z is, independently, H, C_1 - C_8 alkyl, C_1 - C_8 haloalkyl, $\text{C}(=\text{NH})\text{N}(\text{H})\text{R}_u$, $\text{C}(=\text{O})\text{N}(\text{H})\text{R}_u$ or $\text{OC}(=\text{O})\text{N}(\text{H})\text{R}_u$;

R_b , R_g and R_h comprise a ring system having from about 4 to about 7 carbon atoms or
10 having from about 3 to about 6 carbon atoms and 1 or 2 heteroatoms wherein the heteroatoms are selected from oxygen, nitrogen and sulfur and wherein the ring system is aliphatic, unsaturated aliphatic, aromatic, or saturated or unsaturated heterocyclic;

R_j is alkyl or haloalkyl having 1 to about 10 carbon atoms, alkenyl having 2 to about 10 carbon atoms, alkynyl having 2 to about 10 carbon atoms, aryl having 6 to about 14 carbon
15 atoms, $\text{N}(\text{R}_k)(\text{R}_m)$ OR_k , halo, SR_k or CN;

m_a is 1 to about 10;

each m_b is, independently, 0 or 1;

m_c is 0 or an integer from 1 to 10;

m_d is an integer from 1 to 10;

20 m_e is from 0, 1 or 2; and

provided that when m_c is 0, m_d is greater than 1.

Representative substituents groups of Formula I are disclosed in U.S. Serial No. 09/130,973, filed August 7, 1998, entitled "Capped 2'-Oxyethoxy Oligonucleotides."

Representative cyclic substituent groups of Formula II are disclosed in U.S. Serial No. 09/123,108, filed July 27, 1998, entitled "RNA Targeted 2'-Oligomeric compounds that are
25 Conformationally Preorganized".

Particular sugar substituent groups include $\text{O}((\text{CH}_2)_n\text{O})_m\text{CH}_3$, $\text{O}(\text{CH}_2)_n\text{OCH}_3$, $\text{O}(\text{CH}_2)_n\text{NH}_2$, $\text{O}(\text{CH}_2)_n\text{CH}_3$, $\text{O}(\text{CH}_2)_n\text{ONH}_2$, and $\text{O}(\text{CH}_2)_n\text{ON}((\text{CH}_2)_n\text{CH}_3)_2$, where n and m are from 1 to about 10.

30 Representative guanidino substituent groups that are shown in formula III and IV are disclosed in U.S. Serial No. 09/349,040, entitled "Functionalized Oligomers", filed July 7, 1999.

Representative acetamido substituent groups are disclosed in U.S. Patent 6,147,200.

Representative dimethylaminoethoxyethyl substituent groups are disclosed in International Patent Application PCT/US99/17895, entitled "2'-O-Dimethylaminoethoxyethyl-Oligomeric compounds", filed August 6, 1999.

The terms "modified internucleoside linkage" and "modified backbone," or simply
5 "modified linkage" as used herein, refer to modifications or replacement of the naturally occurring phosphodiester internucleoside linkage connecting two adjacent nucleosides within an oligomeric compound. Such modified linkages include those that have a phosphorus atom and those that do not have a phosphorus atom.

Internucleoside linkages containing a phosphorus atom therein include, for example,
10 phosphorothioates, chiral phosphorothioates, phosphorodithioates, phosphotriesters, aminoalkylphosphotriesters, methyl and other alkyl phosphonates including 3'-alkylene phosphonates, 5'-alkylene phosphonates and chiral phosphonates, phosphinates, phosphoramidates including 3'-amino phosphoramidate and aminoalkylphosphoramidates, thionophosphoramidates, thionoalkylphosphonates, thionoalkylphosphotriesters,
15 selenophosphates and boranophosphates having normal 3'-5' linkages, 2'-5' linked analogs of these, and those having inverted polarity wherein one or more internucleotide linkages is a 3' to 3', 5' to 5' or 2' to 2' linkage. Oligonucleotides having inverted polarity can comprise a single 3' to 3' linkage at the 3'-most internucleotide linkage i.e. a single inverted nucleoside residue which may be abasic (the nucleobase is missing or has a hydroxyl group in place thereof). Various
20 salts, mixed salts and free acid forms are also included. Representative U.S. patents that teach the preparation of the above phosphorus-containing linkages include, but are not limited to, U.S.: 3,687,808; 4,469,863; 4,476,301; 5,023,243; 5,177,196; 5,188,897; 5,264,423; 5,276,019; 5,278,302; 5,286,717; 5,321,131; 5,399,676; 5,405,939; 5,453,496; 5,455,233; 5,466,677; 5,476,925; 5,519,126; 5,536,821; 5,541,306; 5,550,111; 5,563,253; 5,571,799; 5,587,361;
25 5,194,599; 5,565,555; 5,527,899; 5,721,218; 5,672,697 and 5,625,050.

In the *C. elegans* system, modification of the internucleotide linkage (phosphorothioate in place of phosphodiester) did not significantly interfere with RNAi activity, indicating that oligomeric compounds of the invention can have one or more modified internucleoside linkages, and retain activity. Indeed, such modified internucleoside linkages are often desired over the
30 naturally occurring phosphodiester linkage because of advantageous properties they can impart such as, for example, enhanced cellular uptake, enhanced affinity for nucleic acid target and increased stability in the presence of nucleases.

Another phosphorus containing modified internucleoside linkage is the phosphonomonoester (see U.S. Patents 5,874,553 and 6,127,346). Phosphonomonoester nucleic acids have

useful physical, biological and pharmacological properties in the areas of inhibiting gene expression (antisense oligonucleotides, ribozymes, sense oligonucleotides and triplex-forming oligonucleotides), as probes for the detection of nucleic acids and as auxiliaries for use in molecular biology.

5 As previously defined an oligonucleoside refers to a sequence of nucleosides that are joined by internucleoside linkages that do not have phosphorus atoms. Non-phosphorus containing internucleoside linkages include short chain alkyl, cycloalkyl, mixed heteroatom alkyl, mixed heteroatom cycloalkyl, one or more short chain heteroatomic and one or more short chain heterocyclic. These internucleoside linkages include but are not limited to siloxane,
10 sulfide, sulfoxide, sulfone, acetyl, formacetyl, thioformacetyl, methylene formacetyl, thioformacetyl, alkenyl, sulfamate; methyleneimino, methylenehydrazino, sulfonate, sulfonamide, amide and others having mixed N, O, S and CH₂ component parts. Representative U.S. patents that teach the preparation of the above oligonucleosides include, but are not limited to, U.S.: 5,034,506; 5,166,315; 5,185,444; 5,214,134; 5,216,141; 5,235,033; 5,264,562;
15 5,264,564; 5,405,938; 5,434,257; 5,466,677; 5,470,967; 5,489,677; 5,541,307; 5,561,225; 5,596,086; 5,602,240; 5,610,289; 5,602,240; 5,608,046; 5,610,289; 5,618,704; 5,623,070; 5,663,312; 5,633,360; 5,677,437; 5,792,608; 5,646,269 and 5,677,439.

Some additional examples of modified internucleoside linkages that do not contain a phosphorus atom therein include, -CH₂-NH-O-CH₂-, -CH₂-N(CH₃)-O-CH₂- (known as a
20 methylene (methylimino) or MMI backbone), -CH₂-O-N(CH₃)-CH₂-, -CH₂-N(CH₃)-N(CH₃)-CH₂- and -O-N(CH₃)-CH₂-CH₂- (wherein the native phosphodiester internucleotide linkage is represented as -O-P(=O)(OH)-O-CH₂-). The MMI type and amide internucleoside linkages are disclosed in the below referenced U.S. patents 5,489,677 and 5,602,240, respectively.

Another modification that can enhance the properties of an oligomeric compound or
25 can be used to track the oligomeric compound or its metabolites is the attachment of one or more moieties or conjugates. Properties that are typically enhanced include without limitation activity, cellular distribution and cellular uptake. In one embodiment, such modified oligomeric compounds are prepared by covalently attaching conjugate groups to functional groups available on an oligomeric compound such as hydroxyl or amino functional groups. Conjugate groups of
30 the invention include intercalators, reporter molecules, polyamines, polyamides, polyethylene glycols, polyethers, groups that enhance the pharmacodynamic properties of oligomers, and groups that enhance the pharmacokinetic properties of oligomers. Typical conjugate groups include cholesterol, lipids, phospholipids, biotin, phenazine, folate, phenanthridine, anthraquinone, acridine, fluoresceins, rhodamines, coumarins, and dyes. Groups that enhance

the pharmacodynamic properties, in the context of this invention, include groups that improve properties including but not limited to oligomer uptake, enhance oligomer resistance to degradation, and/or strengthen sequence-specific hybridization with RNA. Groups that enhance the pharmacokinetic properties, in the context of this invention, include groups that improve
5 properties including but not limited to oligomer uptake, distribution, metabolism and excretion. Representative conjugate groups are disclosed in International Patent Application PCT/US92/09196.

Conjugate groups include but are not limited to lipid moieties such as a cholesterol moiety (Letsinger et al., Proc. Natl. Acad. Sci. USA, 1989, 86, 6553-6556), cholic acid
10 (Manoharan et al., Bioorg. Med. Chem. Lett., 1994, 4, 1053-1060), a thioether, e.g., hexyl-S-tritylthiol (Manoharan et al., Ann. N.Y. Acad. Sci., 1992, 660, 306-309; Manoharan et al., Bioorg. Med. Chem. Lett., 1993, 3, 2765-2770), a thiocholesterol (Oberhauser et al., Nucl. Acids Res., 1992, 20, 533-538), an aliphatic chain, e.g., dodecandiol or undecyl residues (Saison-Behmoaras et al., EMBO J., 1991, 10, 1111-1118; Kabanov et al., FEBS Lett., 1990, 259, 327-
15 330; Svinarchuk et al., Biochimie, 1993, 75, 49-54), a phospholipid, e.g., di-hexadecyl-rac-glycerol or triethylammonium 1,2-di-O-hexadecyl-rac-glycero-3-H-phosphonate (Manoharan et al., Tetrahedron Lett., 1995, 36, 3651-3654; Shea et al., Nucl. Acids Res., 1990, 18, 3777-3783), a polyamine or a polyethylene glycol chain (Manoharan et al., Nucleosides & Nucleotides, 1995, 14, 969-973), or adamantane acetic acid (Manoharan et al., Tetrahedron Lett., 1995, 36, 3651-
20 3654), a palmityl moiety (Mishra et al., Biochim. Biophys. Acta, 1995, 1264, 229-237), or an octadecylamine or hexylamino-carbonyl-oxycholesterol moiety (Crooke et al., J. Pharmacol. Exp. Ther., 1996, 277, 923-937).

The oligomeric compounds of the invention may also be conjugated to active drug substances, for example, aspirin, warfarin, phenylbutazone, ibuprofen, suprofen, fenbufen,
25 ketoprofen, (S)-(+)-pranoprofen, carprofen, dansylsarcosine, 2,3,5-triiodobenzoic acid, flufenamic acid, folic acid, a benzothiadiazide, chlorothiazide, a diazepine, indomethacin, a barbiturate, a cephalosporin, a sulfa drug, an antidiabetic, an antibacterial or an antibiotic. Oligonucleotide-drug conjugates and their preparation are described in U.S. Patent Application 09/334,130.

30 Representative U.S. patents that teach the preparation of such oligonucleotide conjugates include, but are not limited to, U.S.: 4,828,979; 4,948,882; 5,218,105; 5,525,465; 5,541,313; 5,545,730; 5,552,538; 5,578,717; 5,580,731; 5,580,731; 5,591,584; 5,109,124; 5,118,802; 5,138,045; 5,414,077; 5,486,603; 5,512,439; 5,578,718; 5,608,046; 4,587,044; 4,605,735; 4,667,025; 4,762,779; 4,789,737; 4,824,941; 4,835,263; 4,876,335; 4,904,582;

4,958,013; 5,082,830; 5,112,963; 5,214,136; 5,082,830; 5,112,963; 5,214,136; 5,245,022;
5,254,469; 5,258,506; 5,262,536; 5,272,250; 5,292,873; 5,317,098; 5,371,241, 5,391,723;
5,416,203, 5,451,463; 5,510,475; 5,512,667; 5,514,785; 5,565,552; 5,567,810; 5,574,142;
5,585,481; 5,587,371; 5,595,726; 5,597,696; 5,599,923; 5,599,928 and 5,688,941.

5 Oligomeric compounds used in the compositions of the present invention can also be
modified to have one or more stabilizing groups that are generally attached to one or both termini
of oligomeric compounds to enhance properties such as for example nuclease stability. Included
in stabilizing groups are cap structures. The terms "cap structure" or "terminal cap moiety," as
used herein, refer to chemical modifications, which can be attached to one or both of the termini
10 of an oligomeric compound. These terminal modifications protect the oligomeric compounds
having terminal nucleic acid moieties from exonuclease degradation, and can help in delivery
and/or localization within a cell. The cap can be present at the 5'-terminus (5'-cap) or at the 3'-
terminus (3'-cap) or can be present on both termini. In non-limiting examples, the 5'-cap
includes inverted abasic residue (moiety), 4',5'-methylene nucleotide; 1-(beta-D-
15 erythrofuransyl) nucleotide, 4'-thio nucleotide, carbocyclic nucleotide; 1,5-anhydrohexitol
nucleotide; L-nucleotides; alpha-nucleotides; modified base nucleotide; phosphorodithioate
linkage; threo-pentofuransyl nucleotide; acyclic 3',4'-seco nucleotide; acyclic 3,4-
dihydroxybutyl nucleotide; acyclic 3,5-dihydroxypentyl nucleotide, 3'-3'-inverted nucleotide
moiety; 3'-3'-inverted abasic moiety; 3'-2'-inverted nucleotide moiety; 3'-2'-inverted abasic
20 moiety; 1,4-butanediol phosphate; 3'-phosphoramidate; hexylphosphate; aminohexyl phosphate;
3'-phosphate; 3'-phosphorothioate; phosphorodithioate; or bridging or non-bridging
methylphosphonate moiety (for more details see Wincott et al., International PCT publication
No. WO 97/26270).

Particularly suitable 3'-cap structures of the present invention include, for example 4',5'-
25 methylene nucleotide; 1-(beta-D-erythrofuransyl) nucleotide; 4'-thio nucleotide, carbocyclic
nucleotide; 5'-amino-alkyl phosphate; 1,3-diamino-2-propyl phosphate, 3-aminopropyl
phosphate; 6-aminohexyl phosphate; 1,2-aminododecyl phosphate; hydroxypropyl phosphate;
1,5-anhydrohexitol nucleotide; L-nucleotide; alpha-nucleotide; modified base nucleotide;
phosphorodithioate; threo-pentofuransyl nucleotide; acyclic 3',4'-seco nucleotide; 3,4-
30 dihydroxybutyl nucleotide; 3,5-dihydroxypentyl nucleotide, 5'-5'-inverted nucleotide moiety; 5'-
5'-inverted abasic moiety; 5'-phosphoramidate; 5'-phosphorothioate; 1,4-butanediol phosphate;
5'-amino; bridging and/or non-bridging 5'-phosphoramidate, phosphorothioate and/or
phosphorodithioate, bridging or non bridging methylphosphonate and 5'-mercapto moieties (for

more details see Beaucage and Tyer, 1993, Tetrahedron 49, 1925 and Published U.S. Patent Application Publication No. US 2005/0020525 published on January 27, 2005).

Further 3' and 5'-stabilizing groups that can be used to cap one or both ends of an oligomeric compound to impart nuclease stability include those disclosed in WO 03/004602.

5 Oligomerization of modified and unmodified nucleosides is performed according to literature procedures for DNA (Protocols for Oligonucleotides and Analogs, Ed. Agrawal (1993), Humana Press) and/or RNA (Scaringe, Methods (2001), 23, 206-217. Gait et al., Applications of Chemically synthesized RNA in RNA:Protein Interactions, Ed. Smith (1998), 1-36. Gallo et al., Tetrahedron (2001), 57, 5707-5713) synthesis as appropriate. In addition specific protocols for
10 the synthesis of oligomeric compounds of the invention are illustrated in the examples below.

Support bound oligonucleotide synthesis relies on sequential addition of nucleotides to one end of a growing chain. Typically, a first nucleoside (having protecting groups on any exocyclic amine functionalities present) is attached to an appropriate glass bead support and nucleotides bearing the appropriate activated phosphite moiety, i.e. an "activated phosphorous
15 group" (typically nucleotide phosphoramidites, also bearing appropriate protecting groups) are added stepwise to elongate the growing oligonucleotide. Additional methods for solid-phase synthesis may be found in Caruthers U.S. Patents Nos. 4,415,732; 4,458,066; 4,500,707; 4,668,777; 4,973,679; and 5,132,418; and Koster U.S. Patents Nos. 4,725,677 and Re. 34,069.

Oligonucleotides are generally prepared either in solution or on a support medium, e.g.
20 a solid support medium. In general a first synthon (e.g. a monomer, such as a nucleoside) is first attached to a support medium, and the oligonucleotide is then synthesized by sequentially coupling monomers to the support-bound synthon. This iterative elongation eventually results in a final oligomeric compound or other polymer such as a polypeptide. Suitable support medium can be soluble or insoluble, or may possess variable solubility in different solvents to allow the
25 growing support bound polymer to be either in or out of solution as desired. Traditional support medium such as solid support media are for the most part insoluble and are routinely placed in reaction vessels while reagents and solvents react with and/or wash the growing chain until the oligomer has reached the target length, after which it is cleaved from the support and, if necessary further worked up to produce the final polymeric compound. More recent approaches
30 have introduced soluble supports including soluble polymer supports to allow precipitating and dissolving the iteratively synthesized product at desired points in the synthesis (Gravert et al., Chem. Rev., 1997, 97, 489-510).

The term support medium is intended to include all forms of support known to one of ordinary skill in the art for the synthesis of oligomeric compounds and related compounds such

as peptides. Some representative support medium that are amenable to the methods of the present invention include but are not limited to the following: controlled pore glass (CPG); oxalyl-controlled pore glass (see, e.g., Alul, et al., *Nucleic Acids Research* 1991, 19, 1527); silica-containing particles, such as porous glass beads and silica gel such as that formed by the reaction of trichloro-[3-(4-chloromethyl)phenyl]propylsilane and porous glass beads (see Parr and Grohmann, *Angew. Chem. International Ed.* 1972, 11, 314, sold under the trademark "PORASIL E" by Waters Associates, Framingham, Mass., USA); the mono ester of 1,4-dihydroxymethylenbenzene and silica (see Bayer and Jung, *Tetrahedron Lett.*, 1970, 4503, sold under the trademark "BIOPAK" by Waters Associates); TENTAGEL (see, e.g., Wright, et al., *Tetrahedron Letters* 1993, 34, 3373); cross-linked styrene/divinylbenzene copolymer beaded matrix or POROS, a copolymer of polystyrene/divinylbenzene (available from Perceptive Biosystems); soluble support medium, polyethylene glycol PEGs (see Bonora et al., *Organic Process Research & Development*, 2000, 4, 225-231).

The term "linking moiety," as used herein is generally a bi-functional group, covalently binds the ultimate 3'-nucleoside (and thus the nascent oligonucleotide) to the solid support medium during synthesis, but which is cleaved under conditions orthogonal to the conditions under which the 5'-protecting group, and if applicable any 2'-protecting group, are removed. Suitable linking moieties include, but are not limited to, a divalent group such as alkylene, cycloalkylene, arylene, heterocyclyl, heteroarylene, and the other variables are as described above.

Exemplary alkylene linking moieties include, but are not limited to, C₁-C₁₂ alkylene (e.g. methylene, ethylene (e.g. ethyl-1,2-ene), propylene (e.g. propyl-1,2-ene, propyl-1,3-ene), butylene, (e.g. butyl-1,4-ene, 2-methylpropyl-1,3-ene), pentylene, hexylene, heptylene, octylene, decylene, dodecylene), etc. Exemplary cycloalkylene groups include C₃-C₁₂ cycloalkylene groups, such as cyclopropylene, cyclobutylene, cyclopentanyl-1,3-ene, cyclohexyl-1,4-ene, etc. Exemplary arylene linking moieties include, but are not limited to, mono- or bicyclic arylene groups having from 6 to about 14 carbon atoms, e.g. phenyl-1,2-ene, naphthyl-1,6-ene, naphthyl-2,7-ene, anthracenyl, etc. Exemplary heterocyclyl groups within the scope of the invention include mono- or bicyclic aryl groups having from about 4 to about 12 carbon atoms and about 1 to about 4 hetero atoms, such as N, O and S, where the cyclic moieties may be partially dehydrogenated.

Certain heteroaryl groups that may be mentioned as being within the scope of the invention include: pyrrolidinyl, piperidinyl (e.g. 2,5-piperidinyl, 3,5-piperidinyl), piperazinyl, tetrahydrothiophenyl, tetrahydrofuranlyl, tetrahydro quinolinyl, tetrahydro isoquinolinyl,

tetrahydroquinazoliny, tetrahydroquinoxaliny, etc. Exemplary heteroarylene groups include mono- or bicyclic aryl groups having from about 4 to about 12 carbon atoms and about 1 to about 4 hetero atoms, such as N, O and S. Certain heteroaryl groups that may be mentioned as being within the scope of the invention include: pyridylene (e.g. pyridyl-2,5-ene, pyridyl-3,5-ene), pyrimidinyl, thiophenyl, furanyl, quinoliny, isoquinoliny, quinazoliny, quinoxaliny, etc.

Commercially available equipment routinely used for the support medium based synthesis of oligomeric compounds and related compounds is sold by several vendors including, for example, Applied Biosystems (Foster City, CA). Any other means for such synthesis known in the art may additionally or alternatively be employed. Suitable solid phase techniques, including automated synthesis techniques, are described in F. Eckstein (ed.), *Oligonucleotides and Analogues, a Practical Approach*, Oxford University Press, New York (1991).

Although a lot of research has focused on the synthesis of oligoribonucleotides the main RNA synthesis strategies that are presently being used commercially include 5'-O-DMT-2'-O-t-butyldimethylsilyl (TBDMS), 5'-O-DMT-2'-O-[1(2-fluorophenyl)-4-methoxypiperidin-4-yl] (FPMP), 2'-O-[(triisopropylsilyl)oxy]methyl (2'-O-CH₂-O-Si(iPr)₃ (TOM), and the 5'-O-silyl ether-2'-ACE (5'-O-bis(trimethylsiloxy)cyclododecyloxysilyl ether (DOD)-2'-O-bis(2-acetoxyethoxy)methyl (ACE). A current list of some of the major companies currently offering RNA products include Pierce Nucleic Acid Technologies, Dharmacon Research Inc., Ameri Biotechnologies Inc., and Integrated DNA Technologies, Inc. One company, Princeton Separations, is marketing an RNA synthesis activator advertised to reduce coupling times especially with TOM and TBDMS chemistries. Such an activator would also be amenable to the present invention. The primary groups being used for commercial RNA synthesis are:

TBDMS =	5'-O-DMT-2'-O-t-butyldimethylsilyl;
TOM	= 2'-O-[(triisopropylsilyl)oxy]methyl;
25 DOD/ACE	= 5'-O-bis(trimethylsiloxy)cyclododecyloxysilylether- 2'-O-bis(2-acetoxyethoxy)methyl;
FPMP	= 5'-O-DMT-2'-O-[1(2-fluorophenyl)-4-methoxypiperidin-4-yl].

All of the aforementioned RNA synthesis strategies are amenable to the present invention. Strategies that would be a hybrid of the above e.g. using a 5'-protecting group from one strategy with a 2'-O-protecting from another strategy is also amenable to the present invention.

The terms "antisense" or "antisense inhibition" as used herein refer to the hybridization of an oligomeric compound or a portion thereof with a selected target nucleic acid. Multiple antisense mechanisms exist by which oligomeric compounds can be used to modulate gene

expression in mammalian cells. Such antisense inhibition is typically based upon hydrogen bonding-based hybridization of complementary strands or segments such that at least one strand or segment is cleaved, degraded, or otherwise rendered inoperable. In this regard, it is presently suitable to target specific nucleic acid molecules and their functions for such antisense inhibition.

5 The functions of DNA to be interfered with can include replication and transcription. Replication and transcription, for example, can be from an endogenous cellular template, a vector, a plasmid construct or otherwise. The functions of RNA to be interfered with can include functions such as translocation of the RNA to a site of protein translation, translocation of the RNA to sites within the cell which are distant from the site of RNA synthesis, translation of
10 protein from the RNA, splicing of the RNA to yield one or more RNA species, and catalytic activity or complex formation involving the RNA which may be engaged in or facilitated by the RNA.

A commonly exploited antisense mechanism is RNase H-dependent degradation of a targeted RNA. RNase H is a ubiquitously expressed endonuclease that recognizes antisense
15 DNA-RNA heteroduplexes, hydrolyzing the RNA strand. A further antisense mechanism involves the utilization of enzymes that catalyze the cleavage of RNA-RNA duplexes. These reactions are catalyzed by a class of RNase enzymes including but not limited to RNase III and RNase L. The antisense mechanism known as RNA interference (RNAi) is operative on RNA-RNA hybrids and the like. Both RNase H-based antisense (usually using single-stranded
20 compounds) and RNA interference (usually using double-stranded compounds known as siRNAs) are antisense mechanisms, typically resulting in loss of target RNA function.

Optimized siRNA and RNase H-dependent oligomeric compounds behave similarly in terms of potency, maximal effects, specificity and duration of action, and efficiency. Moreover it has been shown that in general, activity of dsRNA constructs correlated with the activity of
25 RNase H-dependent single-stranded antisense oligomeric compounds targeted to the same site. One major exception is that RNase H-dependent antisense oligomeric compounds were generally active against target sites in pre-mRNA whereas siRNAs were not.

These data suggest that, in general, sites on the target RNA that were not active with RNase H-dependent oligonucleotides were similarly not good sites for siRNA. Conversely, a
30 significant degree of correlation between active RNase H oligomeric compounds and siRNA was found, suggesting that if a site is available for hybridization to an RNase H oligomeric compound, then it is also available for hybridization and cleavage by the siRNA complex. Consequently, once suitable target sites have been determined by either antisense approach, these sites can be used to design constructs that operate by the alternative antisense mechanism

(Vickers et al., J. Biol. Chem., 2003, 278, 7108). Moreover, once a site has been demonstrated as active for either an RNAi or an RNase H oligomeric compound, a single-stranded RNAi oligomeric compound (ssRNAi or asRNA) can be designed.

The oligomeric compounds and methods of the present invention are also useful in the study, characterization, validation and modulation of small non-coding RNAs. These include, but are not limited to, microRNAs (miRNA), small nuclear RNAs (snRNA), small nucleolar RNAs (snoRNA), small temporal RNAs (stRNA) and tiny non-coding RNAs (tncRNA) or their precursors or processed transcripts or their association with other cellular components.

Small non-coding RNAs have been shown to function in various developmental and regulatory pathways in a wide range of organisms, including plants, nematodes and mammals. MicroRNAs are small non-coding RNAs that are processed from larger precursors by enzymatic cleavage and inhibit translation of mRNAs. stRNAs, while processed from precursors much like miRNAs, have been shown to be involved in developmental timing regulation. Other non-coding small RNAs are involved in events as diverse as cellular splicing of transcripts, translation, transport, and chromosome organization.

As modulators of small non-coding RNA function, the oligomeric compounds of the present invention find utility in the control and manipulation of cellular functions or processes such as regulation of splicing, chromosome packaging or methylation, control of developmental timing events, increase or decrease of target RNA expression levels depending on the timing of delivery into the specific biological pathway and translational or transcriptional control. In addition, the oligomeric compounds of the present invention can be modified in order to optimize their effects in certain cellular compartments, such as the cytoplasm, nucleus, nucleolus or mitochondria.

The compounds of the present invention can further be used to identify components of regulatory pathways of RNA processing or metabolism as well as in screening assays or devices.

Targeting an oligomeric compound to a particular nucleic acid molecule, in the context of this invention, can be a multistep process. The process usually begins with the identification of a target nucleic acid whose function is to be modulated. The terms "target nucleic acid" and "nucleic acid target", as used herein, refer to any nucleic acid capable of being targeted including without limitation DNA (a cellular gene), RNA (including pre-mRNA and mRNA or portions thereof) transcribed from such DNA, and also cDNA derived from such RNA. In one embodiment the modulation of expression of a selected gene is associated with a particular disorder or disease state. In another embodiment the target nucleic acid is a nucleic acid molecule from an infectious agent.

The targeting process usually also includes determination of at least one target region, segment, or site within the target nucleic acid for the antisense interaction to occur such that the desired effect, e.g., modulation of expression, will result. Within the context of the present invention as it is applied to a nucleic acid target, the term "region" is defined as a portion of the target nucleic acid having at least one identifiable structure, function, or characteristic. Within regions of target nucleic acids are segments. "Segments" are defined as smaller or sub-portions of regions within a target nucleic acid. "Sites," as used in the present invention, are defined as positions within a target nucleic acid. The terms region, segment, and site can also be used to describe an oligomeric compound of the invention such as for example a gapped oligomeric compound having 3 separate regions or segments.

Since, as is known in the art, the translation initiation codon is typically 5'-AUG (in transcribed mRNA molecules; 5'-ATG in the corresponding DNA molecule), the translation initiation codon is also referred to as the "AUG codon," the "start codon" or the "AUG start codon". A minority of genes have a translation initiation codon having the RNA sequence 5'-GUG, 5'-UUG or 5'-CUG, and 5'-AUA, 5'-ACG and 5'-CUG have been shown to function *in vivo*. Thus, the terms "translation initiation codon" and "start codon" can encompass many codon sequences, even though the initiator amino acid in each instance is typically methionine (in eukaryotes) or formylmethionine (in prokaryotes). It is also known in the art that eukaryotic and prokaryotic genes may have two or more alternative start codons, any one of which may be preferentially utilized for translation initiation in a particular cell type or tissue, or under a particular set of conditions. In the context of the invention, "start codon" and "translation initiation codon" refer to the codon or codons that are used *in vivo* to initiate translation of an mRNA transcribed from a gene encoding a nucleic acid target, regardless of the sequence(s) of such codons. It is also known in the art that a translation termination codon (or "stop codon") of a gene may have one of three sequences, i.e., 5'-UAA, 5'-UAG and 5'-UGA (the corresponding DNA sequences are 5'-TAA, 5'-TAG and 5'-TGA, respectively).

The terms "start codon region" and "translation initiation codon region" refer to a portion of such an mRNA or gene that encompasses from about 25 to about 50 contiguous nucleotides in either direction (i.e., 5' or 3') from a translation initiation codon. Similarly, the terms "stop codon region" and "translation termination codon region" refer to a portion of such an mRNA or gene that encompasses from about 25 to about 50 contiguous nucleotides in either direction (i.e., 5' or 3') from a translation termination codon. Consequently, the "start codon region" (or "translation initiation codon region") and the "stop codon region" (or "translation

termination codon region”) are all regions which may be targeted effectively with the antisense oligomeric compounds of the present invention.

The open reading frame (ORF) or “coding region,” which is known in the art to refer to the region between the translation initiation codon and the translation termination codon, is also
5 a region which may be targeted effectively. Within the context of the present invention, one region is the intragenic region encompassing the translation initiation or termination codon of the open reading frame (ORF) of a gene.

Other target regions include the 5' untranslated region (5'UTR), known in the art to refer to the portion of an mRNA in the 5' direction from the translation initiation codon, and thus
10 including nucleotides between the 5' cap site and the translation initiation codon of an mRNA (or corresponding nucleotides on the gene), and the 3' untranslated region (3'UTR), known in the art to refer to the portion of an mRNA in the 3' direction from the translation termination codon, and thus including nucleotides between the translation termination codon and 3' end of an mRNA (or corresponding nucleotides on the gene). The 5' cap site of an mRNA comprises an N7-
15 methylated guanosine residue joined to the 5'-most residue of the mRNA via a 5'-5' triphosphate linkage. The 5' cap region of an mRNA is considered to include the 5' cap structure itself as well as the first 50 nucleotides adjacent to the cap site. It is also suitable to target the 5' cap region.

Although some eukaryotic mRNA transcripts are directly translated, many contain one or more regions, known as “introns,” which are excised from a transcript before it is translated.
20 The remaining (and therefore translated) regions are known as “exons” and are spliced together to form a continuous mRNA sequence. Targeting splice sites, i.e., intron-exon junctions or exon-intron junctions, may also be particularly useful in situations where aberrant splicing is implicated in disease, or where an overproduction of a particular splice product is implicated in disease. Aberrant fusion junctions due to rearrangements or deletions are also suitable target
25 sites. mRNA transcripts produced via the process of splicing of two (or more) mRNAs from different gene sources are known as “fusion transcripts”. It is also known that introns can be effectively targeted using antisense oligomeric compounds targeted to, for example, DNA or pre-mRNA.

It is also known in the art that alternative RNA transcripts can be produced from the
30 same genomic region of DNA. These alternative transcripts are generally known as “variants”. More specifically, “pre-mRNA variants” are transcripts produced from the same genomic DNA that differ from other transcripts produced from the same genomic DNA in either their start or stop position and contain both intronic and exonic sequences.

Upon excision of one or more exon or intron regions, or portions thereof during splicing, pre-mRNA variants produce smaller "mRNA variants". Consequently, mRNA variants are processed pre-mRNA variants and each unique pre-mRNA variant must always produce a unique mRNA variant as a result of splicing. These mRNA variants are also known as
5 "alternative splice variants". If no splicing of the pre-mRNA variant occurs then the pre-mRNA variant is identical to the mRNA variant.

It is also known in the art that variants can be produced through the use of alternative signals to start or stop transcription and that pre-mRNAs and mRNAs can possess more than one start codon or stop codon. Variants that originate from a pre-mRNA or mRNA that use
10 alternative start codons are known as "alternative start variants" of that pre-mRNA or mRNA. Those transcripts that use an alternative stop codon are known as "alternative stop variants" of that pre-mRNA or mRNA. One specific type of alternative stop variant is the "polyA variant" in which the multiple transcripts produced result from the alternative selection of one of the "polyA stop signals" by the transcription machinery, thereby producing transcripts that terminate at
15 unique polyA sites. Within the context of the invention, the types of variants described herein are also suitable target nucleic acids.

The locations on the target nucleic acid to which the antisense oligomeric compounds hybridize are hereinbelow referred to as "suitable target segments." As used herein the term "suitable target segment" is defined as at least an 8-nucleobase portion of a target region to
20 which an active antisense oligomeric compound is targeted. While not wishing to be bound by theory, it is presently believed that these target segments represent portions of the target nucleic acid which are accessible for hybridization.

Exemplary antisense oligomeric compounds include oligomeric compounds that comprise at least the 8 consecutive nucleobases from the 5'-terminus of a targeted nucleic acid
25 e.g. a cellular gene or mRNA transcribed from the gene (the remaining nucleobases being a consecutive stretch of the same oligonucleotide beginning immediately upstream of the 5'-terminus of the antisense oligomeric compound which is specifically hybridizable to the target nucleic acid and continuing until the oligonucleotide contains from about 8 to about 80 nucleobases). Similarly, antisense oligomeric compounds are represented by oligonucleotide
30 sequences that comprise at least the 8 consecutive nucleobases from the 3'-terminus of one of the illustrative antisense oligomeric compounds (the remaining nucleobases being a consecutive stretch of the same oligonucleotide beginning immediately downstream of the 3'-terminus of the antisense oligomeric compound which is specifically hybridizable to the target nucleic acid and continuing until the oligonucleotide contains from about 8 to about 80 nucleobases). One having

skill in the art armed with the antisense oligomeric compounds illustrated herein will be able, without undue experimentation, to identify further antisense oligomeric compounds.

Once one or more target regions, segments or sites have been identified, antisense oligomeric compounds are chosen which are sufficiently complementary to the target, i.e.,
5 hybridize sufficiently well and with sufficient specificity, to give the desired effect.

In accordance with one embodiment of the present invention, a series of nucleic acid duplexes comprising the antisense oligomeric compounds of the present invention and their complements can be designed for a specific target or targets. The ends of the strands may be modified by the addition of one or more natural or modified nucleobases to form an overhang.
10 The sense strand of the duplex is then designed and synthesized as the complement of the antisense strand and may also contain modifications or additions to either terminus. For example, in one embodiment, both strands of the duplex would be complementary over the central nucleobases, each having overhangs at one or both termini.

RNA strands of the duplex can be synthesized by methods disclosed herein or
15 purchased from various RNA synthesis companies such as for example Dharmacon Research Inc., (Lafayette, CO). Once synthesized, the complementary strands are annealed. The single strands are aliquoted and diluted to a concentration of 50 μ M. Once diluted, 30 μ L of each strand is combined with 15 μ L of a 5X solution of annealing buffer. The final concentration of the buffer is 100 mM potassium acetate, 30 mM HEPES-KOH pH 7.4, and 2mM magnesium
20 acetate. The final volume is 75 μ L. This solution is incubated for 1 minute at 90°C and then centrifuged for 15 seconds. The tube is allowed to sit for 1 hour at 37°C at which time the dsRNA duplexes are used in experimentation. The final concentration of the dsRNA compound is 20 μ M. This solution can be stored frozen (-20°C) and freeze-thawed up to 5 times.

Once prepared, the desired synthetic duplexes are evaluated for their ability to modulate
25 target expression. When cells reach 80% confluency, they are treated with synthetic duplexes comprising at least one oligomeric compound of the invention. For cells grown in 96-well plates, wells are washed once with 200 μ L OPTI-MEM-1 reduced-serum medium (Gibco BRL) and then treated with 130 μ L of OPTI-MEM-1 containing 12 μ g/mL LIPOFECTIN (Gibco BRL) and the desired dsRNA compound at a final concentration of 200 nM. After 5 hours of
30 treatment, the medium is replaced with fresh medium. Cells are harvested 16 hours after treatment, at which time RNA is isolated and target reduction measured by RT-PCR.

In a further embodiment, the "suitable target segments" identified herein may be employed in a screen for additional oligomeric compounds that modulate the expression of a target. "Modulators" are those oligomeric compounds that decrease or increase the expression of

a nucleic acid molecule encoding a target and which comprise at least an 8-nucleobase portion which is complementary to a suitable target segment. The screening method comprises the steps of contacting a suitable target segment of a nucleic acid molecule encoding a target with one or more candidate modulators, and selecting for one or more candidate modulators which decrease
5 or increase the expression of a nucleic acid molecule encoding a target. Once it is shown that the candidate modulator or modulators are capable of modulating (e.g. either decreasing or increasing) the expression of a nucleic acid molecule encoding a target, the modulator may then be employed in further investigative studies of the function of a target, or for use as a research, diagnostic, or therapeutic agent in accordance with the present invention.

10 The suitable target segments of the present invention may also be combined with their respective complementary antisense oligomeric compounds of the present invention to form stabilized double stranded (duplexed) oligonucleotides.

In the context of this invention, "hybridization" means hydrogen bonding, which may be Watson-Crick, Hoogsteen or reversed Hoogsteen hydrogen bonding, between the heterocyclic
15 base moieties of complementary nucleosides. For example, adenine and thymine are complementary nucleobases which pair through the formation of hydrogen bonds.

"Complementary," as used herein, refers to the capacity for precise pairing between two nucleotides. For example, if a nucleotide at a certain position of an oligonucleotide is capable of hydrogen bonding with a nucleotide at the same position of a DNA or RNA molecule, then the
20 oligonucleotide and the DNA or RNA are considered to be complementary to each other at that position. The oligonucleotide and the DNA or RNA are complementary to each other when a sufficient number of corresponding positions in each molecule are occupied by nucleotides which can hydrogen bond with each other. Thus, "specifically hybridizable" and "complementary" are terms which are used to indicate a sufficient degree of complementarity or
25 precise pairing such that stable and specific binding occurs between the oligonucleotide and the DNA or RNA target. It is understood in the art that the sequence of an antisense oligomeric compound need not be 100% complementary to that of its target nucleic acid to be specifically hybridizable. An antisense oligomeric compound is specifically hybridizable when binding of the compound to the target DNA or RNA molecule interferes with the normal function of the
30 target DNA or RNA to cause a complete or partial loss of function, and there is a sufficient degree of complementarity to avoid non-specific binding of the antisense oligomeric compound to non-target sequences under conditions in which specific binding is desired, i.e., under physiological conditions in the case of therapeutic treatment, or under conditions in which *in vitro* or *in vivo* assays are performed. Moreover, an oligonucleotide may hybridize over one or

more segments such that intervening or adjacent segments are not involved in the hybridization event (e.g., a loop structure, mismatch or hairpin structure).

The oligomeric compounds of the present invention comprise at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 99%, or 100% sequence complementarity to a target region within the target nucleic acid sequence to which they are targeted. For example, an antisense oligomeric compound in which 18 of 20 nucleobases of the antisense oligomeric compound are complementary to a target region, and would therefore specifically hybridize, would represent 90 percent complementarity. In this example, the remaining noncomplementary nucleobases may be clustered or interspersed with complementary nucleobases and need not be contiguous to each other or to complementary nucleobases. As such, an antisense oligomeric compound which is 18 nucleobases in length having 4 (four) noncomplementary nucleobases which are flanked by two regions of complete complementarity with the target nucleic acid would have 77.8% overall complementarity with the target nucleic acid and would thus fall within the scope of the present invention.

Percent complementarity of an antisense oligomeric compound with a region of a target nucleic acid can be determined routinely using BLAST programs (basic local alignment search tools) and PowerBLAST programs known in the art (Altschul et al., J. Mol. Biol., 1990, 215, 403-410; Zhang and Madden, Genome Res., 1997, 7, 649-656). Percent homology, sequence identity or complementarity, can be determined by, for example, the Gap program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, Madison WI), using default settings, which uses the algorithm of Smith and Waterman (Adv. Appl. Math., 1981, 2, 482-489). In some embodiments, homology, sequence identity or complementarity, between the oligomeric compound and the target is about 70%, about 75%, about 80%, about 85%, about 90%, about 92%, about 94%, about 95%, about 96%, about 97%, about 98%, about 99%, or 100%.

In some embodiments, "suitable target segments" may be employed in a screen for additional oligomeric compounds that modulate the expression of a selected protein.

"Modulators" are those oligomeric compounds that decrease or increase the expression of a nucleic acid molecule encoding a protein and which comprise at least an 8-nucleobase portion which is complementary to a suitable target segment. The screening method comprises the steps of contacting a suitable target segment of a nucleic acid molecule encoding a protein with one or more candidate modulators, and selecting for one or more candidate modulators which decrease or increase the expression of a nucleic acid molecule encoding a protein. Once it is shown that the candidate modulator or modulators are capable of modulating (e.g. either decreasing or

increasing) the expression of a nucleic acid molecule encoding a peptide, the modulator may then be employed in further investigative studies of the function of the peptide, or for use as a research, diagnostic, or therapeutic agent in accordance with the present invention.

The suitable target segments of the present invention may also be combined with their
5 respective complementary antisense oligomeric compounds of the present invention to form stabilized double stranded (duplexed) oligonucleotides. Such double stranded oligonucleotide moieties have been shown in the art to modulate target expression and regulate translation as well as RNA processing via an antisense mechanism. Moreover, the double stranded moieties may be subject to chemical modifications (Fire et al., *Nature*, 1998, 391, 806-811; Timmons and
10 Fire, *Nature* 1998, 395, 854; Timmons et al., *Gene*, 2001, 263, 103-112; Tabara et al., *Science*, 1998, 282, 430-431; Montgomery et al., *Proc. Natl. Acad. Sci. USA*, 1998, 95, 15502-15507; Tuschl et al., *Genes Dev.*, 1999, 13, 3191-3197; Elbashir et al., *Nature*, 2001, 411, 494-498; Elbashir et al., *Genes Dev.* 2001, 15, 188-200). For example, such double stranded moieties have been shown to inhibit the target by the classical hybridization of antisense strand of the
15 duplex to the target, thereby triggering enzymatic degradation of the target (Tijsterman et al., *Science*, 2002, 295, 694-697). The oligomeric compounds of the present invention can also be applied in the areas of drug discovery and target validation. The present invention comprehends the use of the oligomeric compounds and targets identified herein in drug discovery efforts to elucidate relationships that exist between proteins and a disease state, phenotype, or condition.
20 These methods include detecting or modulating a target peptide comprising contacting a sample, tissue, cell, or organism with the oligomeric compounds of the present invention, measuring the nucleic acid or protein level of the target and/or a related phenotypic or chemical endpoint at some time after treatment, and optionally comparing the measured value to a non-treated sample or sample treated with a further oligomeric compound of the invention. These methods can also
25 be performed in parallel or in combination with other experiments to determine the function of unknown genes for the process of target validation or to determine the validity of a particular gene product as a target for treatment or prevention of a particular disease, condition, or phenotype.

Effect of nucleoside modifications on RNAi activity can be evaluated according to
30 existing literature (Elbashir et al., *Nature*, 2001, 411, 494-498; Nishikura et al., *Cell*, 2001, 107, 415-416; and Bass et al., *Cell*, 2000, 101, 235-238.)

The oligomeric compounds of the present invention can be utilized for diagnostics, therapeutics, prophylaxis and as research reagents and kits. Furthermore, antisense oligonucleotides, which are able to inhibit gene expression with exquisite specificity, are often

used by those of ordinary skill to elucidate the function of particular genes or to distinguish between functions of various members of a biological pathway. For use in kits and diagnostics, the oligomeric compounds of the present invention, either alone or in combination with other oligomeric compounds or therapeutics, can be used as tools in differential and/or combinatorial analyses to elucidate expression patterns of a portion or the entire complement of genes expressed within cells and tissues.

As one nonlimiting example, expression patterns within cells or tissues treated with one or more antisense oligomeric compounds are compared to control cells or tissues not treated with antisense oligomeric compounds and the patterns produced are analyzed for differential levels of gene expression as they pertain, for example, to disease association, signaling pathway, cellular localization, expression level, size, structure or function of the genes examined. These analyses can be performed on stimulated or unstimulated cells and in the presence or absence of other compounds and or oligomeric compounds which affect expression patterns.

Examples of methods of gene expression analysis known in the art include DNA arrays or microarrays (Brazma and Vilo, FEBS Lett., 2000, 480, 17-24; Celis, et al., FEBS Lett., 2000, 480, 2-16), SAGE (serial analysis of gene expression)(Madden, et al., Drug Discov. Today, 2000, 5, 415-425), READS (restriction enzyme amplification of digested cDNAs) (Prashar and Weissman, Methods Enzymol., 1999, 303, 258-72), TOGA (total gene expression analysis) (Sutcliffe, et al., Proc. Natl. Acad. Sci. U. S. A., 2000, 97, 1976-81), protein arrays and proteomics (Celis, et al., FEBS Lett., 2000, 480, 2-16; Jungblut, et al., Electrophoresis, 1999, 20, 2100-10), expressed sequence tag (EST) sequencing (Celis, et al., FEBS Lett., 2000, 480, 2-16; Larsson, et al., J. Biotechnol., 2000, 80, 143-57), subtractive RNA fingerprinting (SuRF) (Fuchs, et al., Anal. Biochem., 2000, 286, 91-98; Larson, et al., Cytometry, 2000, 41, 203-208), subtractive cloning, differential display (DD) (Jurecic and Belmont, Curr. Opin. Microbiol., 2000, 3, 316-21), comparative genomic hybridization (Carulli, et al., J. Cell Biochem. Suppl., 1998, 31, 286-96), FISH (fluorescent in situ hybridization) techniques (Going and Gusterson, Eur. J. Cancer, 1999, 35, 1895-904) and mass spectrometry methods (To, Comb. Chem. High Throughput Screen, 2000, 3, 235-41).

The oligomeric compounds of the invention are useful for research and diagnostics, in one aspect because they hybridize to nucleic acids encoding proteins. For example, oligonucleotides that are shown to hybridize with such efficiency and under such conditions as disclosed herein as to be effective protein inhibitors will also be effective primers or probes under conditions favoring gene amplification or detection, respectively. These primers and probes are useful in methods requiring the specific detection of nucleic acid molecules encoding

proteins and in the amplification of the nucleic acid molecules for detection or for use in further studies. Hybridization of the antisense oligonucleotides, particularly the primers and probes, of the invention with a nucleic acid can be detected by means known in the art. Such means may include conjugation of an enzyme to the oligonucleotide, radiolabelling of the oligonucleotide or
5 any other suitable detection means. Kits using such detection means for detecting the level of selected proteins in a sample may also be prepared.

The specificity and sensitivity of antisense is also harnessed by those of skill in the art for therapeutic uses. Antisense oligomeric compounds have been employed as therapeutic moieties in the treatment of disease states in animals, including humans. Antisense
10 oligonucleotide drugs, including ribozymes, have been safely and effectively administered to humans and numerous clinical trials are presently underway. It is thus established that antisense oligomeric compounds can be useful therapeutic modalities that can be configured to be useful in treatment regimes for the treatment of cells, tissues and animals, especially humans.

As used herein, the term "patient" refers to a mammal that is afflicted with one or more
15 disorders associated with expression or overexpression of one or more genes. It will be understood that the most suitable patient is a human. It is also understood that this invention relates specifically to the inhibition of mammalian expression or overexpression of one or more genes.

It is recognized that one skilled in the art may affect the disorders associated with
20 expression or overexpression of a gene by treating a patient presently afflicted with the disorders with an effective amount of one or more oligomeric compounds or compositions of the present invention. Thus, the terms "treatment" and "treating" are intended to refer to all processes wherein there may be a slowing, interrupting, arresting, controlling, or stopping of the progression of the disorders described herein, but does not necessarily indicate a total elimination
25 of all symptoms.

As used herein, the term "effective amount" or "therapeutically effective amount" of a compound of the present invention refers to an amount that is effective in treating or preventing the disorders described herein.

For therapeutics, a patient, such as a human, suspected of having a disease or disorder
30 which can be treated by modulating the expression of a gene is treated by administering antisense oligomeric compounds in accordance with this invention. The compounds of the invention can be utilized in pharmaceutical compositions by adding an effective amount of an antisense oligomeric compound to a suitable pharmaceutically acceptable diluent or carrier. Use of the antisense oligomeric compounds and methods of the invention may also be useful

prophylactically, e.g., to prevent or delay infection, inflammation or tumor formation, for example. In some embodiments, the patient being treated has been identified as being in need of treatment or has been previously diagnosed as such.

The oligomeric compounds of the invention encompass any pharmaceutically acceptable salts, esters, or salts of such esters, or any other compound which, upon administration to an animal including a human, is capable of providing (directly or indirectly) the biologically active metabolite or residue thereof. Accordingly, for example, the disclosure is also drawn to prodrugs and pharmaceutically acceptable salts of the compounds of the invention, pharmaceutically acceptable salts of such prodrugs, and other bioequivalents. For oligonucleotides, examples of pharmaceutically acceptable salts and their uses are further described in U.S. Patent 6,287,860.

The compositions of the invention may also be admixed, encapsulated, conjugated or otherwise associated with other molecules, molecule structures or mixtures of compounds, as for example, liposomes, receptor-targeted molecules, oral, rectal, topical or other formulations, for assisting in uptake, distribution and/or absorption. Representative U.S. patents that teach the preparation of such uptake, distribution and/or absorption-assisting formulations include, but are not limited to, U.S.: 5,108,921; 5,354,844; 5,416,016; 5,459,127; 5,521,291; 5,543,158; 5,547,932; 5,583,020; 5,591,721; 4,426,330; 4,534,899; 5,013,556; 5,108,921; 5,213,804; 5,227,170; 5,264,221; 5,356,633; 5,395,619; 5,416,016; 5,417,978; 5,462,854; 5,469,854; 5,512,295; 5,527,528; 5,534,259; 5,543,152; 5,556,948; 5,580,575; and 5,595,756.

The present invention also includes pharmaceutical compositions and formulations which include the compositions of the invention. The pharmaceutical compositions of the present invention may be administered in a number of ways depending upon whether local or systemic treatment is desired and upon the area to be treated. Administration may be topical (including ophthalmic and to mucous membranes including vaginal and rectal delivery), pulmonary, e.g., by inhalation or insufflation of powders or aerosols, including by nebulizer; intratracheal, intranasal, epidermal and transdermal), oral or parenteral. Parenteral administration includes intravenous, intraarterial, subcutaneous, intraperitoneal or intramuscular injection or infusion; or intracranial, e.g., intrathecal or intraventricular, administration.

Oligonucleotides with at least one 2'-O-methoxyethyl modification are believed to be particularly useful for oral administration. Pharmaceutical compositions and formulations for topical administration may include transdermal patches, ointments, lotions, creams, gels, drops, suppositories, sprays, liquids and powders. Conventional pharmaceutical carriers, aqueous,

powder or oily bases, thickeners and the like may be necessary or desirable. Coated condoms, gloves and the like may also be useful.

Pharmaceutical compositions of the present invention include, but are not limited to, solutions, emulsions, foams and liposome-containing formulations. The pharmaceutical
5 compositions and formulations of the present invention may comprise one or more penetration enhancers, carriers, excipients or other active or inactive ingredients.

One of skill in the art will recognize that formulations are routinely designed according to their intended use, i.e. route of administration.

Suitable formulations for topical administration include those in which the
10 oligonucleotides of the invention are in admixture with a topical delivery agent such as lipids, liposomes, fatty acids, fatty acid esters, steroids, chelating agents and surfactants. Suitable lipids and liposomes include neutral (e.g. dioleoylphosphatidyl DOPE ethanolamine, dimyristoylphosphatidyl choline DMPC, distearoylphosphatidyl choline) negative (e.g. dimyristoylphosphatidyl glycerol DMPG) and cationic (e.g. dioleoyltetramethylaminopropyl
15 DOTAP and dioleoylphosphatidyl ethanolamine DOTMA). Penetration enhancers and their uses are further described in U.S. Patent 6,287,860. Surfactants and their uses are further described in U.S. Patent 6,287,860.

Compositions and formulations for oral administration include powders or granules, microparticulates, nanoparticulates, suspensions or solutions in water or non-aqueous media,
20 capsules, gel capsules, sachets, tablets or minitables. Thickeners, flavoring agents, diluents, emulsifiers, dispersing aids or binders may be desirable. Suitable oral formulations are those in which oligonucleotides of the invention are administered in conjunction with one or more penetration enhancers surfactants and chelators. Suitable surfactants include fatty acids and/or esters or salts thereof, bile acids and/or salts thereof. Suitable bile acids/salts and fatty acids and
25 their uses are further described in U.S. Patent 6,287,860. Also suitable are combinations of penetration enhancers, for example, fatty acids/salts in combination with bile acids/salts. A particularly suitable combination is the sodium salt of lauric acid, capric acid and UDCA. Further penetration enhancers include polyoxyethylene-9-lauryl ether, polyoxyethylene-20-cetyl ether. Oligonucleotides of the invention may be delivered orally, in granular form including
30 sprayed dried particles, or complexed to form micro or nanoparticles. Oligonucleotide complexing agents and their uses are further described in U.S. Patent 6,287,860. Oral formulations for oligonucleotides and their preparation are described in detail in U.S. applications 09/108,673 (filed July 1, 1998), 09/315,298 (filed May 20, 1999) and 10/071,822, filed February 8, 2002.

In another related embodiment, therapeutically effective combination therapies may comprise the use of two or more compositions of the invention wherein the multiple compositions are targeted to a single or multiple nucleic acid targets. Numerous examples of antisense oligomeric compounds are known in the art. Two or more combined compounds may
5 be used together or sequentially.

The formulation of therapeutic compositions and their subsequent administration is believed to be within the skill of those in the art. Dosing is dependent on severity and responsiveness of the disease state to be treated, with the course of treatment lasting from several days to several months, or until a cure is effected or a diminution of the disease state is achieved.
10 Optimal dosing schedules can be calculated from measurements of drug accumulation in the body of the patient. Persons of ordinary skill can easily determine optimum dosages, dosing methodologies and repetition rates. Optimum dosages may vary depending on the relative potency of individual oligonucleotides, and can generally be estimated based on EC₅₀s found to be effective in *in vitro* and *in vivo* animal models. In general, dosage is from 0.01 µg to 100 g
15 per kg of body weight, and may be given once or more daily, weekly, monthly or yearly. Persons of ordinary skill in the art can easily estimate repetition rates for dosing based on measured residence times and concentrations of the drug in bodily fluids or tissues. Following successful treatment, it may be desirable to have the patient undergo maintenance therapy to prevent the recurrence of the disease state, wherein the oligonucleotide is administered in
20 maintenance doses, ranging from 0.01 µg to 100 g per kg of body weight, once or more daily, weekly, monthly, or yearly. For double-stranded compounds, the dose must be calculated to account for the increased nucleic acid load of the second strand (as with compounds comprising two separate strands) or the additional nucleic acid length (as with self complementary single strands having double-stranded regions).

25 While the present invention has been described with specificity in accordance with certain of its embodiments, the following examples serve only to illustrate the invention and are not intended to limit the same.

Examples

30 General

The sequences listed in the examples have been annotated to indicate where there are modified nucleosides or internucleoside linkages. All non-annotated nucleosides are β-D-ribonucleosides linked by phosphodiester internucleoside linkages. Phosphorothioate internucleoside linkages are indicated by underlining. Modified nucleosides are indicated by a

subscripted letter following the capital letter indicating the nucleoside. In particular, subscript "F" indicates 2'-fluoro; subscript "m" indicates 2'-O-methyl; subscript "I" indicates LNA; subscript "e" indicates 2'-O-methoxyethyl (MOE); and subscript "t" indicates 4'-thio. For example U_m is a modified uridine having a 2'-OCH₃ group. A "d" preceding a nucleoside indicates a deoxynucleoside such as dT which is deoxythymidine. Some of the strands have a 5'-phosphate group designated as "P-". Bolded and italicized "***C***" indicates a 5-methyl C ribonucleoside. Where noted next to the ISIS number of a compound, "as" designates the antisense strand, and "s" designates the sense strand of the duplex, with respect to the target sequence.

Example 1: Synthesis of Nucleoside Phosphoramidites

The preparation of nucleoside phosphoramidites is performed following procedures that are extensively illustrated in the art such as but not limited to US Patent 6,426,220 and published PCT WO 02/36743.

Example 2: Oligonucleotide and oligonucleoside synthesis

The oligomeric compounds used in accordance with this invention may be conveniently and routinely made through the well-known technique of solid phase synthesis. Equipment for such synthesis is sold by several vendors including, for example, Applied Biosystems (Foster City, CA). Any other means for such synthesis known in the art may additionally or alternatively be employed. It is well known to use similar techniques to prepare oligonucleotides such as the phosphorothioates and alkylated derivatives.

Oligonucleotides: Unsubstituted and substituted phosphodiester (P=O) oligonucleotides are synthesized on an automated DNA synthesizer (Applied Biosystems model 394) using standard phosphoramidite chemistry with oxidation by iodine.

Phosphorothioates (P=S) are synthesized similar to phosphodiester oligonucleotides with the following exceptions: thiation was effected by utilizing a 10% w/v solution of 3,4-dihydro-2H-benzodithiole-3-one 1,1-dioxide in acetonitrile for the oxidation of the phosphite linkages. The thiation reaction step time was increased to 180 sec and preceded by the normal capping step. After cleavage from the CPG column and deblocking in concentrated ammonium hydroxide at 55°C (12-16 hr), the oligonucleotides were recovered by precipitating with >3 volumes of ethanol from a 1 M NH₄OAc solution. Phosphinate oligonucleotides are prepared as described in U.S. Patent 5,508,270.

Alkyl phosphonate oligonucleotides are prepared as described in U.S. Patent 4,469,863.

3'-Deoxy-3'-methylene phosphonate oligonucleotides are prepared as described in U.S. Patents 5,610,289 or 5,625,050.

Phosphoramidite oligonucleotides are prepared as described in U.S. Patent, 5,256,775 or U.S. Patent 5,366,878.

5 Alkylphosphonothioate oligonucleotides are prepared as described in published PCT applications PCT/US94/00902 and PCT/US93/06976 (published as WO 94/17093 and WO 94/02499, respectively).

3'-Deoxy-3'-amino phosphoramidate oligonucleotides are prepared as described in U.S. Patent 5,476,925.

10 Phosphotriester oligonucleotides are prepared as described in U.S. Patent 5,023,243.

Borano phosphate oligonucleotides are prepared as described in U.S. Patents 5,130,302 and 5,177,198.

Oligonucleosides: Methylenemethylimino linked oligonucleosides, also identified as MMI linked oligonucleosides, methylenedimethylhydrazo linked oligonucleosides, also
15 identified as MDH linked oligonucleosides, and methylenecarbonylamino linked oligonucleosides, also identified as amide-3 linked oligonucleosides, and methyleneaminocarbonyl linked oligonucleosides, also identified as amide-4 linked oligonucleosides, as well as mixed backbone oligomeric compounds having, for instance, alternating MMI and P=O or P=S linkages are prepared as described in U.S. Patents 5,378,825, 5,386,023,
20 5,489,677, 5,602,240 and 5,610,289.

Formacetal and thioformacetal linked oligonucleosides are prepared as described in U.S. Patents 5,264,562 and 5,264,564.

Ethylene oxide linked oligonucleosides are prepared as described in U.S. Patent 5,223,618.

25

Example 3: Oligonucleotide Isolation

After cleavage from the controlled pore glass solid support and deblocking in concentrated ammonium hydroxide at 55°C for 12-16 hours, the oligonucleotides or oligonucleosides are recovered by precipitation out of 1 M NH₄OAc with >3 volumes of ethanol.
30 Synthesized oligonucleotides were analyzed by electrospray mass spectroscopy (molecular weight determination) and by capillary gel electrophoresis and judged to be at least 70% full length material. The relative amounts of phosphorothioate and phosphodiester linkages obtained in the synthesis was determined by the ratio of correct molecular weight relative to the -16 amu product (+/-32 +/-48). For some studies oligonucleotides were purified by HPLC, as described

by Chiang et al., J. Biol. Chem. 1991, 266, 18162-18171. Results obtained with HPLC-purified material were similar to those obtained with non-HPLC purified material.

Example 4: Oligonucleotide Synthesis - 96 Well Plate Format

5 Oligonucleotides can be synthesized via solid phase P(III) phosphoramidite chemistry on an automated synthesizer capable of assembling 96 sequences simultaneously in a 96-well format. Phosphodiester internucleotide linkages are afforded by oxidation with aqueous iodine. Phosphorothioate internucleotide linkages are generated by sulfurization utilizing 3,4-dihydro-2H-benzodithiole-3-one 1,1 dioxide (Beaucage Reagent) in anhydrous acetonitrile. Standard base-
10 protected beta-cyanoethyl-diisopropyl phosphoramidites are purchased from commercial vendors (e.g. PE-Applied Biosystems, Foster City, CA, or Pharmacia, Piscataway, NJ). Non-standard nucleosides are synthesized as per standard or patented methods. They are utilized as base protected beta-cyanoethyldiisopropyl phosphoramidites.

Oligonucleotides are cleaved from support and deprotected with concentrated NH_4OH
15 at elevated temperature (55-60°C) for 12-16 hours and the released product then dried *in vacuo*. The dried product is then re-suspended in sterile water to afford a master plate from which all analytical and test plate samples are then diluted utilizing robotic pipettors.

Example 5: Oligonucleotide Analysis using 96-Well Plate Format

20 The concentration of oligonucleotide in each well is assessed by dilution of samples and UV absorption spectroscopy. The full-length integrity of the individual products is evaluated by capillary electrophoresis (CE) in either the 96-well format (Beckman P/ACE™ MDQ) or, for individually prepared samples, on a commercial CE apparatus (e.g., Beckman P/ACE™ 5000, ABI 270). Base and backbone composition is confirmed by mass analysis of the
25 oligomeric compounds utilizing electrospray-mass spectroscopy. All assay test plates are diluted from the master plate using single and multi-channel robotic pipettors. Plates are judged to be acceptable if at least 85% of the oligomeric compounds on the plate are at least 85% full length.

Example 6: Cell culture and oligonucleotide treatment

30 The effect of oligomeric compounds on target nucleic acid expression can be tested in any of a variety of cell types provided that the target nucleic acid is present at measurable levels. This can be routinely determined using, for example, PCR or Northern blot analysis. Cell lines derived from multiple tissues and species can be obtained from American Type Culture Collection (ATCC, Manassas, VA).

The following cell types are provided for illustrative purposes, but other cell types can be routinely used, provided that the target is expressed in the cell type chosen. This can be readily determined by methods routine in the art, for example Northern blot analysis, ribonuclease protection assays or RT-PCR.

- 5 T-24 cells: The human transitional cell bladder carcinoma cell line T-24 is obtained from the American Type Culture Collection (ATCC) (Manassas, VA). T-24 cells are routinely cultured in complete McCoy's 5A basal media (Invitrogen Corporation, Carlsbad, CA) supplemented with 10% fetal calf serum (Invitrogen Corporation, Carlsbad, CA), penicillin 100 units per mL, and streptomycin 100 micrograms per mL (Invitrogen Corporation, Carlsbad, CA).
10 Cells are routinely passaged by trypsinization and dilution when they reached 90% confluence. Cells are seeded into 96-well plates (Falcon-Primaria #353872) at a density of 7000 cells/well for uses including but not limited to oligomeric compound transfection experiments.

- A549 cells: The human lung carcinoma cell line A549 was obtained from the American Type Culture Collection (Manassas, VA). A549 cells were routinely cultured in
15 DMEM, high glucose (Invitrogen Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum, 100 units per ml penicillin, and 100 micrograms per ml streptomycin (Invitrogen Life Technologies, Carlsbad, CA). Cells were routinely passaged by trypsinization and dilution when they reached approximately 90% confluence. Cells were seeded into 96-well plates (Falcon-Primaria #3872) at a density of approximately 5000 cells/well for uses including
20 but not limited to oligomeric compound transfection experiments.

- b.END cells: The mouse brain endothelial cell line b.END was obtained from Dr. Werner Risau at the Max Plank Institute (Bad Nauheim, Germany). b.END cells were routinely cultured in DMEM, high glucose (Invitrogen Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum (Invitrogen Life Technologies, Carlsbad, CA). Cells were routinely
25 passaged by trypsinization and dilution when they reached approximately 90% confluence. Cells were seeded into 96-well plates (Falcon-Primaria #353872, BD Biosciences, Bedford, MA) at a density of approximately 3000 cells/well for uses including but not limited to oligomeric compound transfection experiments.

- HeLa cells: The human epitheloid carcinoma cell line HeLa was obtained from the
30 American Tissue Type Culture Collection (Manassas, VA). HeLa cells were routinely cultured in DMEM, high glucose (Invitrogen Corporation, Carlsbad, CA) supplemented with 10% fetal bovine serum (Invitrogen Corporation, Carlsbad, CA). Cells were routinely passaged by trypsinization and dilution when they reached 90% confluence. Cells were seeded into 24-well plates (Falcon-Primaria #3846) at a density of 50,000 cells/well or in 96-well plates at a density

of 5,000 cells/well for uses including but not limited to oligomeric compound transfection experiments.

MH-S cells: The mouse alveolar macrophage cell line was obtained from American Type Culture Collection (Manassas, VA). MH-S cells were cultured in RPMI Medium 1640
5 with L-glutamine (Invitrogen Life Technologies, Carlsbad, CA), supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate and 10mM HEPES (all supplements from Invitrogen Life Technologies, Carlsbad, CA). Cells were routinely passaged by trypsinization and dilution when they reached 70-80% confluence. Cells were seeded into 96-well plates (Falcon-Primaria #353047, BD Biosciences, Bedford, MA) at a density of 6500 cells/well for uses including but
10 not limited to oligomeric compound transfection experiments.

U-87 MG: The human glioblastoma U-87 MG cell line was obtained from the American Type Culture Collection (Manassas, VA). U-87 MG cells were cultured in DMEM (Invitrogen Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum (Invitrogen Life Technologies, Carlsbad, CA) and antibiotics. Cells were routinely passaged by
15 trypsinization and dilution when they reached appropriate confluence. Cells were seeded into 96-well plates (Falcon-Primaria #3872) at a density of about 10,000 cells/well for uses including but not limited to oligomeric compound transfection experiments.

Experiments involving treatment of cells with oligomeric compounds:

When cells reach appropriate confluency, they are treated with oligomeric compounds
20 using a transfection method as described.

LIPOFECTIN™

When cells reached 65-75% confluency, they were treated with oligonucleotide. Oligonucleotide was mixed with LIPOFECTIN™ (Invitrogen Life Technologies, Carlsbad, CA) in Opti-MEM™-1 reduced serum medium (Invitrogen Life Technologies, Carlsbad, CA) to
25 achieve the desired concentration of oligonucleotide and a LIPOFECTIN™ concentration of 2.5 or 3 µg/mL per 100 nM oligonucleotide. This transfection mixture was incubated at room temperature for approximately 0.5 hours. For cells grown in 96-well plates, wells were washed once with 100 µL OPTI-MEM™-1 and then treated with 130 µL of the transfection mixture. Cells grown in 24-well plates or other standard tissue culture plates are treated similarly, using
30 appropriate volumes of medium and oligonucleotide. Cells are treated and data are obtained in duplicate or triplicate. After approximately 4-7 hours of treatment at 37°C, the medium containing the transfection mixture was replaced with fresh culture medium. Cells were harvested 16-24 hours after oligonucleotide treatment.

Other suitable transfection reagents known in the art include, but are not limited to, CYTOFECTIN™, LIPOFECTAMINE™, OLIGOFECTAMINE™, and FUGENET™. Other suitable transfection methods known in the art include, but are not limited to, electroporation.

The concentration of oligonucleotide used varies from cell line to cell line. To
 5 determine the optimal oligonucleotide concentration for a particular cell line, the cells are treated with a positive control oligonucleotide at a range of concentrations. For human cells the positive control oligonucleotide is selected from either ISIS 13920
 (T_eC_eC_eGTCATCGCTC_eC_eT_eC_eA_eG_eG_eG_e, SEQ ID NO: 1) which is targeted to human H-ras, or
 ISIS 18078, (G_eT_eG_eC_eG_eCGCGAGCCCG_eA_eA_eA_eT_eC_e, SEQ ID NO: 2) which is targeted to
 10 human Jun-N-terminal kinase-2 (JNK2). Both controls are 2'-O-methoxyethyl gapmers with a phosphorothioate backbone. For mouse or rat cells the positive control oligonucleotide is ISIS
 15770 (A_eT_eG_eC_eA_eTTCTGCCCCCA_eA_eG_eG_eA_e, SEQ ID NO: 3), a 2'-O-methoxyethyl gapmer
 with a phosphorothioate backbone which is targeted to both mouse and rat c-ras. The
 concentration of positive control oligonucleotide that results in 80% inhibition of c-H-ras (for
 15 ISIS 13920), JNK2 (for ISIS 18078) or c-ras (for ISIS 15770) mRNA is then utilized as the
 screening concentration for new oligonucleotides in subsequent experiments for that cell line. If
 80% inhibition is not achieved, the lowest concentration of positive control oligonucleotide that
 results in 60% inhibition of c-H-ras, JNK2 or c-ras mRNA is then utilized as the oligonucleotide
 screening concentration in subsequent experiments for that cell line. If 60% inhibition is not
 20 achieved, that particular cell line is deemed as unsuitable for oligonucleotide transfection
 experiments.

Example 7: Analysis of oligonucleotide inhibition of a target expression

Antisense modulation of a target expression can be assayed in a variety of ways known
 25 in the art. For example, a target mRNA levels can be quantitated by, e.g., Northern blot analysis,
 competitive polymerase chain reaction (PCR), or real-time PCR. Real-time quantitative PCR is
 presently desired. RNA analysis can be performed on total cellular RNA or poly(A)+ mRNA.
 One method of RNA analysis of the present invention is the use of total cellular RNA as
 described in other examples herein. Methods of RNA isolation are well known in the art.
 30 Northern blot analysis is also routine in the art. Real-time quantitative (PCR) can be
 conveniently accomplished using the commercially available ABI PRISM™ 7600, 7700, or 7900
 Sequence Detection System, available from PE-Applied Biosystems, Foster City, CA and used
 according to manufacturer's instructions.

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Protein levels of a target can be quantitated in a variety of ways well known in the art, such as immunoprecipitation, Western blot analysis (immunoblotting), enzyme-linked immunosorbent assay (ELISA) or fluorescence-activated cell sorting (FACS). Antibodies directed to a target can be identified and obtained from a variety of sources, such as the MSRS catalog of antibodies (Aerie Corporation, Birmingham, MI), or can be prepared via conventional monoclonal or polyclonal antibody generation methods well known in the art. Methods for preparation of polyclonal antisera are taught in, for example, Ausubel, F.M. et al., *Current Protocols in Molecular Biology*, Volume 2, pp. 11.12.1-11.12.9, John Wiley & Sons, Inc., 1997. Preparation of monoclonal antibodies is taught in, for example, Ausubel, F.M. et al., *Current Protocols in Molecular Biology*, Volume 2, pp. 11.4.1-11.11.5, John Wiley & Sons, Inc., 1997.

Immunoprecipitation methods are standard in the art and can be found at, for example, Ausubel, F.M. et al., *Current Protocols in Molecular Biology*, Volume 2, pp. 10.16.1-10.16.11, John Wiley & Sons, Inc., 1998. Western blot (immunoblot) analysis is standard in the art and can be found at, for example, Ausubel, F.M. et al., *Current Protocols in Molecular Biology*, Volume 2, pp. 10.8.1-10.8.21, John Wiley & Sons, Inc., 1997. Enzyme-linked immunosorbent assays (ELISA) are standard in the art and can be found at, for example, Ausubel, F.M. et al., *Current Protocols in Molecular Biology*, Volume 2, pp. 11.2.1-11.2.22, John Wiley & Sons, Inc., 1991.

Example 8: Design of phenotypic assays and *in vivo* studies for the use of target inhibitors
Phenotypic assays

Once target inhibitors have been identified by the methods disclosed herein, the oligomeric compounds are further investigated in one or more phenotypic assays, each having measurable endpoints predictive of efficacy in the treatment of a particular disease state or condition.

Phenotypic assays, kits and reagents for their use are well known to those skilled in the art and are herein used to investigate the role and/or association of a target in health and disease. Representative phenotypic assays, which can be purchased from any one of several commercial vendors, include those for determining cell viability, cytotoxicity, proliferation or cell survival (Molecular Probes, Eugene, OR; PerkinElmer, Boston, MA), protein-based assays including enzymatic assays (Panvera, LLC, Madison, WI; BD Biosciences, Franklin Lakes, NJ; Oncogene Research Products, San Diego, CA), cell regulation, signal transduction, inflammation, oxidative processes and apoptosis (Assay Designs Inc., Ann Arbor, MI), triglyceride accumulation (Sigma-Aldrich, St. Louis, MO), angiogenesis assays, tube formation assays, cytokine and hormone

assays and metabolic assays (Chemicon International Inc., Temecula, CA; Amersham Biosciences, Piscataway, NJ).

In one non-limiting example, cells determined to be appropriate for a particular phenotypic assay (i.e., MCF-7 cells selected for breast cancer studies; adipocytes for obesity studies) are treated with a target inhibitors identified from the *in vitro* studies as well as control compounds at optimal concentrations which are determined by the methods described above. At the end of the treatment period, treated and untreated cells are analyzed by one or more methods specific for the assay to determine phenotypic outcomes and endpoints.

Phenotypic endpoints include changes in cell morphology over time or treatment dose as well as changes in levels of cellular components such as proteins, lipids, nucleic acids, hormones, saccharides or metals. Measurements of cellular status which include pH, stage of the cell cycle, intake or excretion of biological indicators by the cell, are also endpoints of interest.

Measurement of the expression of one or more of the genes of the cell after treatment is also used as an indicator of the efficacy or potency of the a target inhibitors. Hallmark genes, or those genes suspected to be associated with a specific disease state, condition, or phenotype, are measured in both treated and untreated cells.

In vivo studies

The individual subjects of the *in vivo* studies described herein are warm-blooded vertebrate animals, which includes humans.

A clinical trial is subjected to rigorous controls to ensure that individuals are not unnecessarily put at risk and that they are fully informed about their role in the study.

To account for the psychological effects of receiving treatments, volunteers are randomly given placebo or a target inhibitor. Furthermore, to prevent the doctors from being biased in treatments, they are not informed as to whether the medication they are administering is a a target inhibitor or a placebo. Using this randomization approach, each volunteer has the same chance of being given either the new treatment or the placebo.

Volunteers receive either the a target inhibitor or placebo for eight week period with biological parameters associated with the indicated disease state or condition being measured at the beginning (baseline measurements before any treatment), end (after the final treatment), and at regular intervals during the study period. Such measurements include the levels of nucleic acid molecules encoding a target or a target protein levels in body fluids, tissues or organs compared to pre-treatment levels. Other measurements include, but are not limited to, indices of the disease state or condition being treated, body weight, blood pressure, serum titers of

pharmacologic indicators of disease or toxicity as well as ADME (absorption, distribution, metabolism and excretion) measurements.

Information recorded for each patient includes age (years), gender, height (cm), family history of disease state or condition (yes/no), motivation rating (some/moderate/great) and
5 number and type of previous treatment regimens for the indicated disease or condition.

Volunteers taking part in this study are healthy adults (age 18 to 65 years) and roughly an equal number of males and females participate in the study. Volunteers with certain characteristics are equally distributed for placebo and a target inhibitor treatment. In general, the volunteers treated with placebo have little or no response to treatment, whereas the volunteers
10 treated with the target inhibitor show positive trends in their disease state or condition index at the conclusion of the study.

Example 9 : RNA Isolation

Poly(A)+ mRNA isolation

15 Poly(A)+ mRNA is isolated according to Miura et al., (Clin. Chem., 1996, 42, 1758-1764). Other methods for poly(A)+ mRNA isolation are routine in the art. Briefly, for cells grown on 96-well plates, growth medium was removed from the cells and each well was washed with 200 μ L cold PBS. 60 μ L lysis buffer (10 mM Tris-HCl, pH 7.6, 1 mM EDTA, 0.5 M NaCl, 0.5% NP-40, 20 mM vanadyl-ribonucleoside complex) was added to each well, the plate was
20 gently agitated and then incubated at room temperature for five minutes. 55 μ L of lysate was transferred to Oligo d(T) coated 96-well plates (AGCT Inc., Irvine CA). Plates were incubated for 60 minutes at room temperature, washed 3 times with 200 μ L of wash buffer (10 mM Tris-HCl pH 7.6, 1 mM EDTA, 0.3 M NaCl). After the final wash, the plate was blotted on paper towels to remove excess wash buffer and then air-dried for 5 minutes. 60 μ L of elution buffer (5
25 mM Tris-HCl pH 7.6), preheated to 70°C, was added to each well, the plate was incubated on a 90°C hot plate for 5 minutes, and the eluate was then transferred to a fresh 96-well plate.

Cells grown on 100 mm or other standard plates may be treated similarly, using appropriate volumes of all solutions.

Total RNA Isolation

30 Total RNA is isolated using an RNEASY 96™ kit and buffers purchased from Qiagen Inc. (Valencia, CA) following the manufacturer's recommended procedures. Briefly, for cells grown on 96-well plates, growth medium is removed from the cells and each well is washed with 200 μ L cold PBS. 150 μ L Buffer RLT is added to each well and the plate vigorously agitated for

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20 seconds. 150 μ L of 70% ethanol is then added to each well and the contents mixed by pipetting three times up and down. The samples are then transferred to the RNEASY 96™ well plate attached to a QIAVAC™ manifold fitted with a waste collection tray and attached to a vacuum source. Vacuum is applied for 1 minute. 500 μ L of Buffer RW1 is added to each well of the RNEASY 96™ plate and incubated for 15 minutes and the vacuum is again applied for 1 minute. An additional 500 μ L of Buffer RW1 is added to each well of the RNEASY 96™ plate and the vacuum is applied for 2 minutes. 1 mL of Buffer RPE is then added to each well of the RNEASY 96™ plate and the vacuum applied for a period of 90 seconds. The Buffer RPE wash is then repeated and the vacuum is applied for an additional 3 minutes. The plate is then removed from the QIAVAC™ manifold and blotted dry on paper towels. The plate is then re-attached to the QIAVAC™ manifold fitted with a collection tube rack containing 1.2 mL collection tubes. RNA is then eluted by pipetting 140 μ L of RNase free water into each well, incubating 1 minute, and then applying the vacuum for 3 minutes.

The repetitive pipetting and elution steps may be automated using a QIAGEN Bio-Robot 9604 (Qiagen, Inc., Valencia CA). Essentially, after lysing of the cells on the culture plate, the plate is transferred to the robot deck where the pipetting, DNase treatment and elution steps are carried out.

Example 10: Design and screening of duplexed antisense compounds

In accordance with the present invention, a series of nucleic acid duplexes comprising the compounds of the present invention and their complements can be designed. The nucleobase sequence of the antisense strand of the duplex comprises at least a portion of an antisense oligonucleotide targeted to a target sequence as described herein. The ends of the strands may be modified by the addition of one or more natural or modified nucleobases to form an overhang.

The sense strand of the dsRNA is then designed and synthesized as the complement of the antisense strand and may also contain modifications or additions to either terminus. For example, in one embodiment, both strands of the dsRNA duplex would be complementary over the central nucleobases, each having overhangs at one or both termini.

For example, a duplex comprising an antisense strand having the sequence CGAGAGGCGGACGGGACCG (SEQ ID NO: 20) and having a two-nucleobase overhang of deoxythymidine(dT) would have the following structure:

cgagaggcggacgggaccgdTdT	Antisense Strand	SEQ ID NO: 21
dTdTgctctccgctgccctggc	Complement Strand	SEQ ID NO: 22

In another embodiment, a duplex comprising an antisense strand having the same sequence CGAGAGGCGGACGGGACCG (SEQ ID NO: 20) may be prepared with blunt ends (no single stranded overhang) as shown:

	cgagaggcggacgggaccg	Antisense Strand SEQ ID NO: 20
5		
	gctctccgcctgccctggc	Complement Strand SEQ ID NO: 23

RNA strands of the duplex can be synthesized by methods disclosed herein or purchased from Dharmacon Research Inc., (Lafayette, CO). Once synthesized, the complementary strands are annealed. The single strands are aliquoted and diluted to a concentration of 50 μ M. Once diluted, 30 μ L of each strand is combined with 15 μ L of a 5X solution of annealing buffer. The final concentration of the buffer is 100 mM potassium acetate, 30 mM HEPES-KOH pH 7.4, and 2mM magnesium acetate. The final volume is 75 μ L. This solution is incubated for 1 minute at 90°C and then centrifuged for 15 seconds. The tube is allowed to sit for 1 hour at 37°C at which time the dsRNA duplexes are used in experimentation.

The final concentration of the dsRNA duplex is 20 μ M.

Once prepared, the duplexed compounds are evaluated for their ability to modulate target mRNA levels. When cells reach 80% confluency, they are treated with duplexed compounds of the invention. For cells grown in 96-well plates, wells are washed once with 200 μ L OPTI-MEM-1™ reduced-serum medium (Gibco BRL) and then treated with 130 μ L of OPTI-MEM-1™ containing 5 μ g/mL LIPOFECTAMINE 2000™ (Invitrogen Life Technologies, Carlsbad, CA) and the duplex antisense compound at the desired final concentration. After about 4 hours of treatment, the medium is replaced with fresh medium. Cells are harvested 16 hours after treatment, at which time RNA is isolated and target reduction measured by quantitative real-time PCR as described herein.

Example 11: Real-time Quantitative PCR Analysis of target mRNA Levels

Quantitation of a target mRNA levels was accomplished by real-time quantitative PCR using the ABI PRISM™ 7600, 7700, or 7900 Sequence Detection System (PE-Applied Biosystems, Foster City, CA) according to manufacturer's instructions. This is a closed-tube, non-gel-based, fluorescence detection system which allows high-throughput quantitation of polymerase chain reaction (PCR) products in real-time. As opposed to standard PCR in which amplification products are quantitated after the PCR is completed, products in real-time quantitative PCR are quantitated as they accumulate. This is accomplished by including in the PCR reaction an oligonucleotide probe that anneals specifically between the forward and reverse

PCR primers, and contains two fluorescent dyes. A reporter dye (e.g., FAM or JOE, obtained from either PE-Applied Biosystems, Foster City, CA, Operon Technologies Inc., Alameda, CA or Integrated DNA Technologies Inc., Coralville, IA) is attached to the 5' end of the probe and a quencher dye (e.g., TAMRA, obtained from either PE-Applied Biosystems, Foster City, CA, Operon Technologies Inc., Alameda, CA or Integrated DNA Technologies Inc., Coralville, IA) is attached to the 3' end of the probe. When the probe and dyes are intact, reporter dye emission is quenched by the proximity of the 3' quencher dye. During amplification, annealing of the probe to the target sequence creates a substrate that can be cleaved by the 5'-exonuclease activity of Taq polymerase. During the extension phase of the PCR amplification cycle, cleavage of the probe by Taq polymerase releases the reporter dye from the remainder of the probe (and hence from the quencher moiety) and a sequence-specific fluorescent signal is generated. With each cycle, additional reporter dye molecules are cleaved from their respective probes, and the fluorescence intensity is monitored at regular intervals by laser optics built into the ABI PRISM™ Sequence Detection System. In each assay, a series of parallel reactions containing serial dilutions of mRNA from untreated control samples generates a standard curve that is used to quantitate the percent inhibition after antisense oligonucleotide treatment of test samples.

Prior to quantitative PCR analysis, primer-probe sets specific to the target gene being measured are evaluated for their ability to be “multiplexed” with a GAPDH amplification reaction. In multiplexing, both the target gene and the internal standard gene GAPDH are amplified concurrently in a single sample. In this analysis, mRNA isolated from untreated cells is serially diluted. Each dilution is amplified in the presence of primer-probe sets specific for GAPDH only, target gene only (“single-plexing”), or both (multiplexing). Following PCR amplification, standard curves of GAPDH and target mRNA signal as a function of dilution are generated from both the single-plexed and multiplexed samples. If both the slope and correlation coefficient of the GAPDH and target signals generated from the multiplexed samples fall within 10% of their corresponding values generated from the single-plexed samples, the primer-probe set specific for that target is deemed multiplexable. Other methods of PCR are also known in the art.

RT and PCR reagents were obtained from Invitrogen Life Technologies (Carlsbad, CA). RT, real-time PCR was carried out by adding 20 µL PCR cocktail (2.5x PCR buffer minus MgCl₂, 6.6 mM MgCl₂, 375 µM each of dATP, dCTP, dGTP and dTTP, 375 nM each of forward primer and reverse primer, 125 nM of probe, 4 Units RNase inhibitor, 1.25 Units PLATINUM® Taq, 5 Units MuLV reverse transcriptase, and 2.5x ROX dye) to 96-well plates containing 30 µL

total RNA solution (20-200 ng). The RT reaction was carried out by incubation for 30 minutes at 48°C. Following a 10 minute incubation at 95°C to activate the PLATINUM® Taq, 40 cycles of a two-step PCR protocol were carried out: 95°C for 15 seconds (denaturation) followed by 60°C for 1.5 minutes (annealing/extension).

5 Gene target quantities obtained by RT, real-time PCR are normalized using either the expression level of GAPDH, a gene whose expression is constant, or by quantifying total RNA using RIBOGREEN™ (Molecular Probes, Inc. Eugene, OR). GAPDH expression is quantified by real time RT-PCR, by being run simultaneously with the target, multiplexing, or separately. Total RNA is quantified using RiboGreen™ RNA quantification reagent (Molecular Probes, Inc. Eugene, OR). Methods of RNA quantification by RIBOGREEN™ are taught in Jones, L.J., et al, (Analytical Biochemistry, 1998, 265, 368-374).

10 In this assay, 170 µL of RIBOGREEN™ working reagent (RIBOGREEN™ reagent diluted 1:350 in 10mM Tris-HCl, 1 mM EDTA, pH 7.5) is pipetted into a 96-well plate containing 30 µL purified, cellular RNA. The plate is read in a CytoFluor 4000 (PE Applied Biosystems) with excitation at 485nm and emission at 530nm.

Example 12: Target-specific primers and probes

Probes and primers may be designed to hybridize to a target sequence, using published sequence information.

20 For example, for human PTEN, the following primer-probe set was designed using published sequence information (GENBANK™ accession number U92436.1, SEQ ID NO: 4).

Forward primer: AATGGCTAAGTGAAGATGACAATCAT (SEQ ID NO: 5)

Reverse primer: TGCACATATCATTACACCAGTTCGT (SEQ ID NO: 6)

And the PCR probe:

25 FAM-TTGCAGCAATTCACTGTAAAGCTGGAAAGG-TAMRA (SEQ ID NO: 7), where FAM is the fluorescent dye and TAMRA is the quencher dye.

For example, for human survivin, the following primer-probe set was designed using published sequence information (GENBANK™ accession number NM_001168.1, SEQ ID NO: 8).

30 Forward primer: CACCACTTCCAGGGTTTATTCC (SEQ ID NO: 9)

Reverse primer: TGATCTCCTTTCCTAAGACATTGCT (SEQ ID NO: 10)

And the PCR probe:

FAM-ACCAGCCTTCCTGTGGGCCCT-TAMRA (SEQ ID NO: 11),

where FAM is the fluorescent dye and TAMRA is the quencher dye.

For example, for human eIF4E, the following primer-probe set was designed using published sequence information (GENBANK™ accession number M15353.1, SEQ ID NO: 12).

Forward primer: TGGCGACTGTCGAACCG (SEQ ID NO: 13)

5 Reverse primer: AGATTCCGTTTTCTCCTCTTCTGTAG (SEQ ID NO: 14)

And the PCR probe:

FAM-AAACCACCCCTACTCCTAATCCCCCG-TAMRA (SEQ ID NO: 15),

where FAM is the fluorescent dye and TAMRA is the quencher dye.

For example, for mouse eIF4E, the following primer-probe set was designed using
10 published sequence information (GENBANK™ accession number NM_007917.2, SEQ ID NO: 16).

Forward primer: AGGACGGTGGCTGATCACA (SEQ ID NO: 17)

Reverse primer: TCTCTAGCCAGAAGCGATCGA (SEQ ID NO: 18)

And the PCR probe:

15 FAM-TGAACAAGCAGCAGAGACGGAGTGA-TAMRA (SEQ ID NO: 19),

where FAM is the fluorescent dye and TAMRA is the quencher dye.

Example 13: Northern blot analysis of a target mRNA levels

Eighteen hours after antisense treatment, cell monolayers were washed twice with cold
20 PBS and lysed in 1 mL RNAZOL™ (TEL-TEST “B” Inc., Friendswood, TX). Total RNA was prepared following manufacturer's recommended protocols. Twenty micrograms of total RNA was fractionated by electrophoresis through 1.2% agarose gels containing 1.1% formaldehyde using a MOPS buffer system (AMRESCO, Inc. Solon, OH). RNA was transferred from the gel to HYBOND™-N+ nylon membranes (Amersham Pharmacia Biotech, Piscataway, NJ) by
25 overnight capillary transfer using a Northern/Southern Transfer buffer system (TEL-TEST “B” Inc., Friendswood, TX). RNA transfer was confirmed by UV visualization. Membranes were fixed by UV cross-linking using a STRATALINKER™ UV Crosslinker 2400 (Stratagene, Inc, La Jolla, CA) and then probed using QUICKHYB™ hybridization solution (Stratagene, La Jolla, CA) using manufacturer's recommendations for stringent conditions.

30 To detect human a target, a human a target specific primer probe set is prepared by PCR. To normalize for variations in loading and transfer efficiency membranes are stripped and probed for human glyceraldehyde-3-phosphate dehydrogenase (GAPDH) RNA (Clontech, Palo Alto, CA).

Hybridized membranes were visualized and quantitated using a PHOSPHORIMAGER™ and IMAGEQUANT™ Software V3.3 (Molecular Dynamics, Sunnyvale, CA). Data was normalized to GAPDH levels in untreated controls.

5 Example 14: Western blot analysis of target protein levels

Western blot analysis (immunoblot analysis) is carried out using standard methods. Cells are harvested 16-20 h after oligonucleotide treatment, washed once with PBS, suspended in Laemmli buffer (100 µl/well), boiled for 5 minutes and loaded on a 16% SDS-PAGE gel. Gels are run for 1.5 hours at 150 V, and transferred to membrane for western blotting. Appropriate
 10 primary antibody directed to a target is used, with a radiolabeled or fluorescently labeled secondary antibody directed against the primary antibody species. Bands are visualized using a PHOSPHORIMAGER™ (Molecular Dynamics, Sunnyvale CA).

Example 15: In vitro assay of selected differentially modified siRNAs

15 Differentially modified siRNA duplexes designed to target human survivin using published sequence information were prepared and assayed as described below. The antisense strand was held constant as a 4'-thio gapped strand and 3 different sense strands were compared. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

20	SEQ ID NO. /ISIS NO.	Composition (5' 3')	Features
	24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	4'-S wings (3/13/3)
	25/352512 (s)	G _m G _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A _m	2'-OCH ₃ full
25	25/352513 (s)	GG _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A	2'-OCH ₃ block (1/17/1)
	25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A	MOE alternating

30 The differentially modified siRNA duplexes were assayed for their ability to inhibit target mRNA levels in HeLa cells. Culture methods used for HeLa cells are available from the ATCC and may be found, for example, at [www\(dot\)atcc.org](http://www(dot)atcc.org). For cells grown in 96-well plates, wells were washed once with 200 µL OPTI-MEM-1 reduced-serum medium and then treated with 130 µL of OPTI-MEM-1 containing 12 µg/mL LIPOFECTIN™ (Invitrogen Life Technologies, Carlsbad, CA) and the dsRNA at the desired concentrations. After about 5 hours

of treatment, the medium was replaced with fresh medium. Cells were harvested 16 hours after treatment, at which time RNA was isolated and target reduction measured by RT-PCR as previously described. Dose-response data was used to determine the IC₅₀ for each pair noted below (antisense:sense).

5	Construct	Assay/Species	Target	IC ₅₀ (nM)
	353537:352512	Dose Response/Human	Survivin	0.60192
	353537:352513	Dose Response/Human	Survivin	0.71193
	353537:352514	Dose Response/Human	Survivin	0.48819.

10 **Example 16: In vitro assay of differentially modified siRNAs having MOE modified sense and 4'-thio (4'-thio/2'-OCH₃) gapmer antisense strands**

In accordance with the present invention, a series of oligomeric compounds were synthesized and tested for their ability to reduce target expression over a range of doses relative to an unmodified compound. The compounds tested were 19 nucleotides in length having

15 phosphorothioate internucleoside linkages throughout.

HeLa cells were treated with the double stranded oligomeric compounds (siRNA constructs) shown below (antisense strand followed by the sense strand of the duplex) at concentrations of 0, 0.15, 1.5, 15, and 150 nM using methods described herein. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

20 Expression levels of human PTEN were determined by quantitative real-time PCR and normalized to RIBOGREEN™ as described in other examples herein. Resulting dose-response curves were used to determine the IC₅₀ for each pair. Also shown is the effect of each duplex on target mRNA levels as a percentage of untreated control (%UTC).

SEQ ID NO.	Composition (5' to 3')	IC ₅₀	%UTC
25 /ISIS NO.			
26/xxxxxx (as)	<u>UUGUCUCUGGUCCUUACUU</u>	0.94	13
27/xxxxxx (s)	<u>AAGUAAGGACCAGAGACAA</u>		
26/xxxxxx (as)	<u>UUGUCUCUGGUCCUUACUU</u>	.055	13
27/359351 (s)	<u>A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e</u>		
30 26/359347 (as)	<u>U_tU_tGUCUCUGGUCCUUACU_tU_t</u>	2.2	25
27/359551 (s)	<u>A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e</u>		
26/359346 (as)	<u>U_tU_tGUCUCUGGUCCUUAC_mU_mU_m</u>	0.18	11
27/359351 (s)	<u>A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e</u>		
26/359345 (as)	<u>U_tU_tGUCUCUGGUCCUUACU_tU_t</u>	5.3	18

27/xxxxxx (s)	<u>AAGUAAGGACCAGAGACAA</u>		
26/359346 (as)	<u>U_tU_tGUCUCUGGUCCUUAC_mU_mU_m</u>	0.73	15
27/xxxxxx (s)	<u>AAGUAAGGACCAGAGACAA</u>		
26/359345 (as)	<u>U_tU_tGUCUCUGGUCCUUACU_tU_t</u>	0.49	14
5 27/xxxxxx (s)	<u>AA_eGU_eAA_eGG_eAC_eCA_eGA_eGA_eCA_eA</u>		
26/359345 (as)	<u>U_tU_tGUCUCUGGUCCUUACU_tU_t</u>	0.55	15
27/359351 (s)	<u>A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e</u>		

From these data it is evident that the activity of the double strand construct containing the 4'-thio gapmer RNA in the antisense strand paired with an RNA sense strand

- 10 (359345_341401 having an IC50 of 5.3) can be improved by incorporating 2'MOE modifications into the sense strand on the terminal ends or in an alternating configuration with RNA. It is also evident that improvements in IC50 values can be obtained over the unmodified pure RNA construct (341391_341401; RNA in both strands with an IC50 value of 0.94) by using an alternating motif.

15

Example 17: In vitro assay of selected differentially modified siRNAs

- Selected siRNAs (shown below as antisense strand followed by the sense strand of the duplex) were prepared and evaluated in HeLa cells treated as described herein with varying doses of the selected siRNAs. The mRNA levels were quantitated using real-time PCR as
- 20 described herein and were compared to untreated control levels (%UTC). The IC50's were calculated using the linear regression equation generated by plotting the normalized mRNA levels to the log of the concentrations used.

SEQ ID NO.	Composition (5' to 3')	IC50	%UTC
/ISIS NO.			
25 26/359346 (as)	U _t U _t GUCUCUGGUCCUUAC _m U _m U _m	1.9	10
27/367287 (s)	AAGU _t AAGGAC _t C _t AGAGAC _t AA		
26/359345 (as)	U _t U _t GUCUCUGGUCCUUACU _t U _t	1.7	20
27/367287 (s)	AAGU _t AAGGAC _t C _t AGAGAC _t AA		
26/359345 (as)	U _t U _t GUCUCUGGUCCUUACU _t U _t	0.2	10
30 27/367288 (s)	A _t A _t GUAAGGACCAGAGACA _t A _t		
26/359346 (as)	U _t U _t GUCUCUGGUCCUUAC _m U _m U _m	< 0.1	10
27/367288 (s)	A _t A _t GUAAGGACCAGAGACA _t A _t		
26/359345 (as)	U _t U _t GUCUCUGGUCCUUACU _t U _t	0.5	15
27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		

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	26/359346 (as)U _t U _t GUCUCUGGUCCUUAAC _m U _m U _m	0.2	11
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/359995 (as)U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m	0.4	17
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
5	26/359345 (as)U _t U _t GUCUCUGGUCCUUAACU _t U _t	0.2	13
	27/359996 (s) A _m A _f G _m U _f A _m A _f G _m G _f A _m C _f C _m A _f G _m A _f G _m A _f C _m A _f A _m		
	26/359346 (as)U _t U _t GUCUCUGGUCCUUAAC _m U _m U _m	0.2	13
	27/359996 (s) A _m A _f G _m U _f A _m A _f G _m G _f A _m C _f C _m A _f G _m A _f G _m A _f C _m A _f A _m		
	26/361203 (as)UUG _m UCUCU _m GGUCC _m UUACU _m U	<0.1	--
10	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/361209 (as)UUGU _m CUCUG _m GUCCU _m UACU _m U	1.5	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/361204 (as)UUGU _e CUCUGG _e UCCUUAACU _e U	1.5	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
15	26/361205 (as)UUGUC _e UCUGGUC _e CUUAC _e U _e U _e	2.5	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/361206 (as)UUGUC _e U _e CUGGU _e C _e CUUACU _e U _e	--	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/361207 (as)UUGUCU _e C _e UGG _e U _e CCUUAAC _e U _e U _e	10.1	--
20	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/341391 (as)UUGUCUCUGGUCCUUAACUU	0.1	--
	27/341401 (s) AAGUAAGGACCAGAGACAA		
	26/359979 (as)UUGUC _m UCU _m GGU _m CCU _m UAC _m U _m U _m	--	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
25	26/359980 (as)UUGUCU _m C _m UGG _m U _m CCUUAAC _m U _m U _m	0.2	--
	27/359351 (s) A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e		
	26/359980 (as)UUGUCU _m C _m UGG _m U _m CCUUAAC _m U _m U _m	0.1	--
	27/361221 (s) A _m A _m G _m UAAGGACCAGAGAC _m A _m A _m		

30 Example 18: In vitro assay of modified siRNAs targeted to human survivin

In accordance with the present invention, a series of oligomeric compounds were synthesized and tested for their ability to reduce survivin expression over a range of doses. HeLa cells were treated with the double stranded oligomeric compounds (siRNA constructs) shown below (antisense strand followed by the sense strand of the duplex) at concentrations of 0.0006

nM, 0.084 nM, 0.16 nM, 0.8 nM, 4 nM, or 20 nM using methods described herein. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. Expression levels of human survivin were determined using real-time PCR methods as described herein. The effect of the 20 nM dose on survivin mRNA levels is shown below.

5 Results are presented as a percentage of untreated control mRNA levels.

SEQ ID NO. Composition (5' to 3')		%UTC
/ISIS NO.		
24/343867 (as)	UUUGAAAAUGUUGAUCUCC	3
25/343868 (s)	GGAGAUCACAUAUUUCAA	
10 24/352506 (as)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m	2
25/371314 (s)	G _e G _e A _e G _e A _e UCAACAUUUU _e C _e A _e A _e	
24/352506 (as)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m	3
25/371316 (s)	G _m G _m A _m GAUCAACAUAUUUCA _m A _m A _m	
24/352506 (as)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m	2
15 25/371313 (s)	G _e G _e A _e GAUCAACAUAUUUCA _e A _e A _e	
24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	5
25/371313 (s)	G _e G _e A _e GAUCAACAUAUUUCA _e A _e A _e	
24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	5
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A	
20 24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	6
25/371314 (s)	G _e G _e A _e G _e A _e UCAACAUAUUUU _e C _e A _e A _e	
24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	5
25/371315 (s)	G _e G _e A _e GAUCAAC _e A _e UUUUCA _e A _e A _e	
24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t	5
25 25/371316 (s)	G _m G _m A _m GAUCAACAUAUUUCA _m A _m A _m	
24/353540 (as)	U _m U _m U _m GAAAAUGUUGAUCU _t C _t C _t	3
25/371313 (s)	G _e G _e A _e GAUCAACAUAUUUCA _e A _e A _e	
24/353540 (as)	U _m U _m U _m GAAAAUGUUGAUCU _t C _t C _t	2
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A	
30 24/353540 (as)	U _m U _m U _m GAAAAUGUUGAUCU _t C _t C _t	3
25/371314 (s)	G _e G _e A _e G _e A _e UCAACAUAUUUU _e C _e A _e A _e	
24/353540 (as)	U _m U _m U _m GAAAAUGUUGAUCU _t C _t C _t	3
25/371315 (s)	G _e G _e A _e GAUCAAC _e A _e UUUUCA _e A _e A _e	
24/353540 (as)	U _m U _m U _m GAAAAUGUUGAUCU _t C _t C _t	3

25/371316 (s)	G _m G _m A _m GAUCAACAUUUUC A _m A _m A _m	
24/368679 (as)	U _m U _f U _m G _f A _m A _f A _m A _f U _m G _f U _m U _f G _m A _f U _m C _f U _m C _f C _m	2
25/371313 (s)	G _e G _e A _e GAUCAACAUUUUC A _e A _e A _e	
24/368679 (as)	U _m U _f U _m G _f A _m A _f A _m A _f U _m G _f U _m U _f G _m A _f U _m C _f U _m C _f C _m	3
5 25/371314 (s)	G _e G _e A _e G _e A _e UCAACAUUUUC C _e A _e A _e A _e	
24/368679 (as)	U _m U _f U _m G _f A _m A _f A _m A _f U _m G _f U _m U _f G _m A _f U _m C _f U _m C _f C _m	3
25/371316 (s)	G _m G _m A _m GAUCAACAUUUUC A _m A _m A _m	
24/352506 (as)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m	12
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A	
10 24/368679 (as)	U _m U _f U _m G _f A _m A _f A _m A _f U _m G _f U _m U _f G _m A _f U _m C _f U _m C _f C _m	8
25/371315 (s)	G _e G _e A _e GAUCAAC _e A _e UUUUC A _e A _e A _e	

Example 19: In vitro assay of selected differentially modified siRNAs targeted to human eIF4E

- 15 In accordance with the present invention, a series of oligomeric compounds were synthesized and tested for their ability to reduce eIF4E expression over a range of doses. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. HeLa cells were treated with the double stranded oligomeric compounds (siRNA constructs) shown below (antisense strand followed by the sense strand to which it was
- 20 duplexed) at concentrations of 0.0006 nM, 0.032 nM, 0.16 nM, 0.8 nM, 4 nM, or 20 nM using methods described herein. Expression levels of human eIF4E were determined using real-time PCR methods as described herein. Resulting dose-response curves were used to determine the IC₅₀ for each pair as shown below.

SEQ ID NO.	Composition (5' to 3')	IC ₅₀
25 /ISIS NO.		
30/371286 (as)	UUUAGCUCUAACAUAACA	0.440
31/371280 (s)	UGUUA AUGUUAGAGCUAAA	
30/371287 (as)	UUUAGC _m U _m CUA _m A _m CAUUA _m C _m A _m	0.356
31/371280 (s)	UGUUA AUGUUAGAGCUAAA	
30 30/371287 (as)	UUUAGC _m U _m CUA _m A _m CAUUA _m C _m A _m	2.520
31/371284 (s)	U _e G _e U _e UAAUGUUAGAGCUA _e A _e A _e	
32/371297 (as)	UUACUAGACAACUGGAU AU	0.381
33/371291 (s)	AUAUCCAGUUGUCUAGUAA	
32/371298 (as)	UUACUA _m G _m ACA _m A _m CUGGAU _m A _m U _m	0.260

33/371291 (s)	AUAUCCAGUUGUCUAGUAA	
32/371298 (as)	UUACUA _m G _m ACA _m A _m CUGGAU _m A _m U _m	0.260
33/371295 (s)	A _e U _e A _e UCCAGUUGUCUAGU _e A _e A _e	
32/379960 (as)	U _m U _f A _m C _f U _m A _f G _m A _f C _m A _f A _m C _f U _m G _f G _m A _f U _m A _f U _m	0.260
5 33/371295 (s)	A _e U _e A _e UCCAGUUGUCUAGU _e A _e A _e	
34/371308 (as)	UUAAAAAGUGAGUAGUCAC	0.126
35/371302 (s)	GUGACUACUCACUUUUUAA	
34/371309 (as)	UUAAAA _m A _m GUG _m A _m GUAGUC _m A _m C _m	0.168
35/371302 (s)	GUGACUACUCACUUUUUAA	
10 34/371309 (as)	UUAAAA _m A _m GUG _m A _m GUAGUC _m A _m C _m	0.040
35/371306 (s)	G _e U _e G _e ACUACUCACUUUUU _e A _e A _e	
34/371309 (as)	UUAAAA _m A _m GUG _m A _m GUAGUC _m A _m C _m	0.017
35/379965 (s)	G _m U _f G _m A _f C _m U _f A _m C _f U _m C _f A _m C _f U _m U _f U _m U _f U _m A _f A _m	

15 **Example 20: In vitro assay of selected differentially modified siRNAs targeted to mouse eIF4E**

In accordance with the present invention, a series of oligomeric compounds were synthesized and tested for their ability to reduce eIF4E expression over a range of doses. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. b.END cells were treated with the double stranded oligomeric compounds (siRNA constructs) shown below (antisense strand followed by the sense strand of the duplex) at concentrations of 0.0625 nM, 0.25 nM, 1 nM, or 4 nM using methods described herein. Expression levels of mouse eIF4E were determined using real-time PCR methods as described herein. Resulting dose-response curves were used to determine the IC₅₀ for each pair as shown below.

SEQ ID NO.	Composition (5' to 3')	IC ₅₀
/ISIS NO.		
30/371286 (as)	UUUAGCUCUAACAUUAACA	0.2055
31/371280 (s)	UGUUA AUGUUAGAGCUAAA	
30 30/371287 (as)	UUUAGC _m U _m CUA _m A _m CAUUA _m C _m A _m	0.238
31/371280 (s)	UGUUA AUGUUAGAGCUAAA	
30/371287 (as)	UUUAGC _m U _m CUA _m A _m CAUUA _m C _m A _m	9.496
31/371284 (s)	U _e G _e U _e UAAUGUUAGAGCUA _e A _e A _e	
30/371286 (as)	UUUAGCUCUAACAUUAACA	1.193

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31/371284 (s)	U _e G _e U _e UAAUGUUAGAGCUA _e A _e A _e	
32/371297 (as)	UUACUAGACAACUGGAUUAU	0.1859
33/371291 (s)	AUAUCCAGUUGUCUAGUAA	
32/371298 (as)	UUACUA _m G _m ACA _m A _m CUGGAU _m A _m U _m	0.1946
5 33/371291 (s)	AUAUCCAGUUGUCUAGUAA	
32/371297 (as)	UUACUAGACAACUGGAUUAU	0.0936
33/371295 (s)	A _e U _e A _e UCCAGUUGUCUAGU _e A _e A _e	
32/371298 (as)	UUACUA _m G _m ACA _m A _m CUGGAU _m A _m U _m	0.1151
33/371295 (s)	A _e U _e A _e UCCAGUUGUCUAGU _e A _e A _e	
10 34/371308 (as)	UUAAAAAGUGAGUAGUCAC	0.2926
35/371302 (s)	GUGACUACUCACUUUUUAA	
34/371309 (as)	UUAAAA _m A _m GUG _m A _m GUAGUC _m A _m C _m	0.1626
35/371302 (s)	GUGACUACUCACUUUUUAA	
34/371308 (as)	UUAAAAAGUGAGUAGUCAC	0.0632
15 35/371306 (s)	G _e U _e G _e ACUACUCACUUUUU _e A _e A _e	
34/371309 (as)	UUAAAA _m A _m GUG _m A _m GUAGUC _m A _m C _m	0.0061
35/371306 (s)	G _e U _e G _e ACUACUCACUUUUU _e A _e A _e	

Example 21: Blockmer walk of 5 2'-O-methy modified nucleosides in the antisense strand of siRNAs assayed for PTEN mRNA levels against untreated control

The antisense (AS) strands listed below were designed to target human PTEN, and each was duplexed with the same sense strand (ISIS 271790, shown below). The duplexes were tested for their ability to reduce PTEN expression over a range of doses to determine the relative positional effect of the 5 modifications using methods described herein. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. Expression levels of PTEN were determined using real-time PCR methods as described herein, and were compared to levels determined for untreated controls.

SEQ ID NO:/ISIS NO	Sequence 5'-3'
36/271790 (S)	CAAAUCCAGAGGCUAGCAGdTdT
30 37/271071(AS)	C _m U _m G _m C _m U _m AGCCUCUGGAUUUGdTdT
37/271072(AS)	CU _m G _m C _m U _m A _m GCCUCUGGAUUUGdTdT
37/271073(AS)	CUG _m C _m U _m A _m G _m CCUCUGGAUUUGdTdT
37/271074(AS)	CUGC _m U _m A _m G _m C _m CUCUGGAUUUGdTdT
37/271075(AS)	CUGC _m U _m A _m G _m C _m C _m UCUGGAUUUGdTdT

The siRNAs having 2'-O-methyl groups at least 2 positions removed from the siRNAs having 5, 2'-O-methyl groups at least 2 positions removed from the 5'-end of the antisense strand reduced PTEN mRNA levels to from 25 to 35% of untreated control. The remaining 2 constructs increased PTEN mRNA levels above untreated control.

5

Example 22: Solid block of 2'-O-methyl modified nucleosides in the antisense strand of siRNAs assayed for PTEN mRNA levels against untreated control

The antisense (AS) strands listed below were designed to target human PTEN, and each was duplexed with the same sense strand 271790. The duplexes were tested for their ability to
 10 reduce PTEN expression over a range of doses to determine the relative effect of adding either 9 or 14, 2'-O-methyl modified nucleosides at the 3'-end of the resulting siRNAs. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. Expression levels of PTEN were determined using real-time PCR methods as described herein, and were compared to levels determined for untreated controls.

15	SEQ ID NO:/ISIS NO	Sequence 5'-3'
	36/271790 (S)	CAAAUCCAGAGGCUAGCAGdTdT
	37/271079(AS)	CUGCUAGCCUCUG _m G _m A _m U _m U _m G _m U _m U _m
	37/271081(AS)	CUGCUAGC _m C _m U _m C _m U _m G _m G _m A _m U _m U _m U _m G _m U _m U _m

The siRNA having 9, 2'-O-methyl nucleosides reduced PTEN mRNA levels to about 40% of
 20 untreated control whereas the construct having 14, 2'-O-methyl nucleosides only reduced PTEN mRNA levels to about 98% of control.

Example 23: 2'-O-methyl blockmers (siRNA vs asRNA)

A series of blockmers were prepared as single strand antisense RNAs (asRNAs). The
 25 antisense (AS) strands listed below were designed to target PTEN, and each was also assayed as part of a duplex with the same sense strand (ISIS 308746, shown below) for their ability to reduce PTEN expression levels. T24 cells were treated with the single stranded or double stranded oligomeric compounds created with the antisense compounds shown below using methods described herein. The nucleosides are annotated as to chemical modification as per the
 30 legend at the beginning of the examples. Expression levels of human PTEN were determined using real-time PCR methods as described herein, and were compared to levels determined for untreated controls.

	SEQ ID NO:/ISIS NO	Sequence 5'-3'
	39/308746 (S)	AAGUAAGGACCAGAGACAAA

40/303912 (AS)	P- <u>UUUGUCUCUGGUCCUUACUU</u>
40/316449 (AS)	P- <u>UUUGUCUCUGGUCCUUAC_mU_mU_m</u>
40/335223 (AS)	P- <u>UUUGUCUCUGGUCCU_mU_mA_mCUU</u>
40/335224 (AS)	P- <u>UUUGUCUCUGGU_mC_mC_mUUACUU</u>
5 40/335225 (AS)	P- <u>UUUGUCUCU_mG_mG_mUCCUUACUU</u>
40/335226 (AS)	P- <u>UUUGUC_mU_mC_mUGGUCCUUACUU</u>
40/335227 (AS)	P- <u>UUU_mG_mU_mCUCUGGUCCUUACUU</u>
40/335228 (AS)	P- <u>U_mU_mU_mGUCUCUGGUCCUUACUU</u>

All of the asRNAs and siRNAs showed activity with the asRNAs having better activity
 10 than the corresponding duplex in each case. A clear dose response was seen for all of the siRNA
 constructs (20, 40, 80 and 150 nm doses). A dose-responsive effect was also observed for the
 asRNAs for 50, 100 and 200 nm doses. In general the siRNAs were more active in this system at
 lower doses than the asRNAs and at the 150 nm dose were able to reduce PTEN mRNA levels to
 15 mRNA levels to about 19% of the untreated control. The duplex containing unmodified 303912 reduced PTEN

Example 24: siRNA hemimer constructs

Three siRNA hemimer constructs were prepared and were tested for their ability to
 reduce PTEN expression levels. The hemimer constructs had 7, 2'-O-methyl nucleosides at the
 20 3'-end. The hemimer was put in the sense strand only, the antisense strand only and in both
 strands to compare the effects. Cells were treated with the double stranded oligomeric
 compounds (siRNA constructs) shown below (antisense strand followed by the sense strand of
 the duplex) using methods described herein. The nucleosides are annotated as to chemical
 modification as per the legend at the beginning of the examples. Expression levels of PTEN
 25 were determined using real-time PCR methods as described herein, and were compared to levels
 determined for untreated controls.

SEQ ID NO:/ISIS NO	Constructs (overhangs) 5'-3'
38/XXXXXX (AS)	<u>CUGCUAGCCUCUGGA_mU_mU_mU_mG_mU_mU_m</u>
41/271068 (S)	CAAAUCCAGAGGCUA _m G _m C _m A _m G _m U _m U _m
30 38/XXXXXX (AS)	<u>CUGCUAGCCUCUGGAUUUGUU</u>
41/271068 (S)	CAAAUCCAGAGGCUA _m G _m C _m A _m G _m U _m U _m
38/XXXXXX (AS)	<u>CUGCUAGCCUCUGGA_mU_mU_mU_mG_mU_mU_m</u>
41/XXXXXX (S)	CAAAUCCAGAGGCUAGCAGUU

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The construct having the 7, 2'-O-methyl nucleosides only in the antisense strand reduced PTEN mRNA levels to about 23% of untreated control. The construct having the 7, 2'-O-methyl nucleosides in both strands reduced the PTEN mRNA levels to about 25% of untreated control. When the 7, 2'-O-methyl nucleosides were only in the sense strand, PTEN mRNA levels were reduced to about 31% of untreated control.

Example 25: Representative siRNAs prepared having 2'-O-Me gapmers

The following antisense strands of selected siRNA duplexes targeting PTEN are hybridized to their complementary full phosphodiester sense strands. Activity is measured using methods described herein. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

SEQ ID NO:	Sequence (5'-3')
42/300852	CUGC _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
42/300853	P-CUGC _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
15 42/300854	C _m U _m G _m C _m UAGCCUCUGGAUU _m U _m G _m A _m
42/300855	P-CUGC _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
42/300856	C _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
42/300858	CUGC _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
42/300859	P-CUGC _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
20 42/300860	C _m U _m A _m G _m CCUCUGGAUU _m U _m G _m A _m
43/303913	G _m U _m C _m U _m CUGGUCCUUA _m C _m U _m U _m
44/303915	U _m U _m U _m U _m GUCUCUGGUC _m C _m U _m U _m
45/303917	C _m U _m G _m G _m UCCUUACUUC _m C _m C _m C _m
46/308743	P-U _m U _m U _m GUCUCUGGUCCUUA _m C _m U _m U _m
25 47/308744	P-U _m C _m U _m C _m U _m GGUCCUUA _m C _m C _m C _m C _m
46/328795	P-UUUG _m U _m C _m U _m CUGGUCCUUA _m C _m U _m U _m

Example 26: Representative siRNAs prepared having 2'-F modified nucleosides and various structural motifs

The following antisense strands of siRNAs targeting PTEN were tested as single strands alone or were hybridized to their complementary full phosphodiester sense strand and were tested in duplex. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. Bolded and italicized "C" indicates a 5-methyl C ribonucleoside.

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SEQ ID NO/ISIS NO		Sequences 5'-3'
40/319022	AS	<u>U_fU_fU_fG_fU_fC_fU_fC_fU_fG_fG_fU_fC_fC_fU_fA_fC_fU_fU_f</u>
40/333749	AS	<u>UUUGUCUCUGGUCCU_fU_fA_fCUU</u>
40/333750	AS	<u>UUUGUCUCUGGU_fC_fC_fUUACUU</u>
5 40/333751	AS	<u>UUUGUCUCUGGU_fC_fC_fUUACUU</u>
40/333752	AS	<u>UUUGUC_fU_fC_fUGGUCCUUACUU</u>
40/333753	AS	<u>UUU_fG_fU_fCUCUGGUCCUUACUU</u>
40/333754	AS	<u>U_fU_fU_fGUCUCUGGUCCUUACUU</u>
40/333756	AS	<u>UUUGUCUCUGGUCCUUAC_fU_fU_f</u>
10 40/334253	AS	<u>UUUGUCUCU_fG_fG_fUCCUUACUU</u>
40/334254	AS	<u>UUUGUCUCUGGUCCUU_fA_fC_fU_fU_f</u>
40/334255	AS	<u>UUU_fG_fU_fCUCUGGUCCUUACUU</u>
40/334256	AS	<u>UUU_fG_fU_fCUCUGGU_fC_fC_fUUACUU</u>
40/334257	AS	<u>U_fU_fU_fGUCUCUGGUCCUUACUU</u>
15 40/317466	AS	<u>U_fU_fU_fGUCUCUGGUCCUUAC_fU_fU_f</u>
40/317468	AS	<u>U_fU_fU_fGUCUCUGGUCCUUAC_fU_fU_f</u>
40/317502	AS	<u>U_fU_fU_fGU_fC_fU_fCUGGUCC_fU_fU_fAC_fU_fU_f</u>

Cells were treated with the indicated concentrations of single or double stranded oligomeric compounds shown above using methods described herein. Expression levels of

- 20 PTEN were determined using real-time PCR methods as described herein, and were compared to levels determined for untreated controls.

% untreated control mRNA		
Construct	100 nM asRNA	100 nM siRNA
303912 35		18
25 317466 --		28
317408 --		18
317502 --		21
334254 --		33
333756 42		19
30 334257 34		23
334255 44		21
333752 42		18
334253 38		15
333750 43		21

333749 34

21

Additional siRNAs having 2'-F modified nucleosides are listed below.

37/279471	AS	$C_fU_fG_fC_fU_fA_fG_fC_fC_fU_fC_fU_fG_fG_fA_fU_fU_fU_fG_dTdT$
5 36/279467	S	$C_fA_fA_fA_fU_fC_fC_fA_fG_fA_fG_fG_fC_fU_fA_fG_fC_fA_fG_dTdT$
40/319018	AS	$U_fU_fU_fG_fU_fC_fU_fC_fU_fG_fG_fU_fC_fC_fU_fU_fA_fC_fU_fU_f$
39/319019	S	$A_fA_fG_fU_fA_fA_fG_fG_fA_fC_fC_fA_fG_fA_fG_fA_fC_fA_fA_fA_f$

Example 27: Representative siRNAs prepared with fully modified antisense strands

10 (2'-F and 2'-OMe)

siRNA constructs targeting PTEN are prepared wherein the following sense and antisense strands are hybridized. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

SEQ ID NO/ISIS NO		Sequences 5'-3'
15 48/283546	(as)	$C_fU_fG_mC_fU_fA_mG_mC_fC_fU_fC_fU_fG_mG_mA_mU_fU_fU_fG_mU_mdT$
40/336240	(s)	<u>UUUGUCUCC_fU_fGGUC_fCUUACmU_mU_m</u>

Example 28: Representative siRNAs prepared having 2'-MOE modified nucleosides were assayed for PTEN mRNA levels against untreated control

20 siRNA constructs targeting PTEN were prepared wherein the following antisense strands were hybridized to the complementary full phosphodiester sense strand.

The following antisense strands of siRNAs were hybridized to the complementary full phosphodiester sense strand. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. Linkages are phosphorothioate. Cells were treated with the duplexes using methods described herein. Results obtained using 100nM duplex are presented as a percentage of untreated control PTEN mRNA levels.

SEQ ID NO. /ISIS NO.	Composition (5' to 3')	PTEN mRNA level (% UTC) 100 nM
49/xxxxxx (as)	<u>UUCAUUCCUGGUCUCUGUUU</u>	--
30 49/xxxxxx (as)	<u>UeU_eCeAUUCCUGGUCUCUGUUU</u>	50
49/xxxxxx (as)	<u>UUCAeU_eUeCCUGGUCUCUGUUU</u>	--
49/xxxxxx (as)	<u>UUCAUUCeC_eUeGGUCUCUGUUU</u>	43
49/xxxxxx (as)	<u>UUCAUUCCUGeG_eUeCUCUGUUU</u>	42
49/xxxxxx (as)	<u>UUCAUUCCUGGUCeU_eCeUGUUU</u>	47

49/xxxxx (as)	<u>UUCAUCCUGGUCUCU_eG_eU_eUU</u>	63
49/xxxxx (as)	<u>UUCAUCCUGGUCUCUGU_eU_eU_e</u>	106

Example 29: 4'-Thio and 2'-OCH₃ chimeric oligomeric compounds

- 5 The double-stranded constructs shown below were prepared (antisense strand followed by the sense strand of the duplex). The "P" following the designation for antisense (as) indicates that the target is PTEN and the "S" indicates that the target is Survivin. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

SEQ ID NO.	Composition (5' to 3')
10 /ISIS NO.	
40/308743 (as-P)	<u>U_mU_mU_mGUCUCUGGUCCUUAC_mU_mU_m</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
24/353537 (as-S)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t
25/343868 (s-S)	GGAGAUCAACAUUUUCAAA
15 24/353537 (as-S)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t
25/352512 (s)	G _m G _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A _m
24/353537 (as-S)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t
25/352513 (s)	GG _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A
24/353537 (as-S)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t
20 25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A

- The constructs designed to the targets indicated were tested in accordance with the assays described herein. The duplexed oligomeric compounds were evaluated in HeLa cells (American Type Culture Collection, Manassas VA). Culture methods used for HeLa cells are available from the ATCC and may be found, for example, at <http://www.atcc.org>. For cells
- 25 grown in 96-well plates, wells were washed once with 200 μ L OPTI-MEM-1 reduced-serum medium and then treated with 130 μ L of OPTI-MEM-1 containing 12 μ g/mL LIPOFECTINTM (Invitrogen Life Technologies, Carlsbad, CA) and the dsRNA at the desired concentration. After about 5 hours of treatment, the medium was replaced with fresh medium. Cells were harvested
- 16 hours after dsRNA treatment, at which time RNA was isolated and target reduction measured
- 30 by quantitative real-time PCR as described in previous examples. Resulting dose-response data was used to determine the IC₅₀ for each construct.

Construct	Assay/Species	Target	IC ₅₀ (nM)
308743:308746	Dose Response/Human	PTEN	0.0275
353537:343868	Dose Response/Human	Survivin	0.067284

353537:343868	Dose Response/Human	Survivin	0.17776
353537:343868	Dose Response/Human	Survivin	0.598
353537:343868	Dose Response/Human	Survivin	4.23
353537:352512	Dose Response/Human	Survivin	0.60192
5 353537:352513	Dose Response/Human	Survivin	0.71193
353537:352514	Dose Response/Human	Survivin	0.48819

Example 30: Selected siRNA constructs prepared and tested against eIF4E and Survivin targets

- 10 Selected siRNA constructs were prepared and tested for their ability to lower targeted RNA as measured by quantitative real-time PCR. The duplexes are shown below (antisense strand followed by the sense strand of the duplex). The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

SEQ ID NO. Composition (5' to 3') Targeted to eIF4E

15 **/ISIS NO.**

50/349894 (as)	U _f G _f U _f C _f A _f UAUCCUGGAU _m C _m C _m U _m U _m
51/338935 (s)	AAGGAUCCAGGAAUAUGACA
52/349895 (as)	U _f C _f C _f U _f G _f GAUCCUUCACC _m A _m A _m U _m G _m
53/338939 (s)	CAUUGGUGAAGGAUCCAGGA
20 54/349896 (as)	U _f C _f U _f U _f A _f UCACCUUUAGC _m U _m C _m U _m A _m
55/338943 (s)	UAGAGCUAAAGGUGAUAAGA
56/349897 (as)	A _f U _f A _f C _f U _f CAGAAGGUGUC _m U _m U _m C _m U _m
57/338952 (s)	AGAAGACACCUUCUGAGUAU
58/352827 (as)	U _s C _s U _s UAUCACCUUUAGCU _m C _m U _m
25 59/342764 (s)	AGAGCUAAAGGUGAUAAGA
58/354604 (as)	U _s C _s U _s U _f A _f U _f C _f A _f C _f C _f U _f U _f A _f G _f C _f U _m C _m U _m
59/342764 (s)	AGAGCUAAAGGUGAUAAGA

SEQ ID NO. Composition (5' to 3') Targeted to Survivin

/ISIS NO.

30 24/355710 (as)	U _f U _f U _f G _f A _f AAAUGUUGAU _m C _m U _m C _m C _m
25/343868 (s)	GGAGAUCAACAUUUUCAAA
24/353540 (as)	U _s U _s U _s GAAAAUGUUGAUCU _m C _m C _m
45/343868 (s)	GGAGAUCAACAUUUUCAAA

The above constructs were tested in HeLa cells, MH-S cells or U-87 MG cells using transfection procedures and real-time PCR as described herein. The resulting IC₅₀'s for the duplexes were calculated and are shown below.

	Construct	Species/cell line	Gene	IC₅₀
5	349894:338935	Human/HeLa	eIF4E	0.165
	349895:338939	Human/HeLa	eIF4E	0.655
	349896:338943	Human/HeLa	eIF4E	0.277
	349896:338943	Mouse/MH-S	eIF4E	0.05771
	349897:338952	Human/HeLa	eIF4E	0.471
10	352827:342764	Human/HeLa	eIF4E	2.033
	352827:342764	Mouse/MH-S	eIF4E	0.34081
	354604:342764	Human/HeLa	eIF4E	2.5765
	355710:343868	Human/HeLa	Survivin	0.048717
	353540:343868	Human/HeLa	Survivin	0.11276
15	353540:343868	Human/U-87 MG	Survivin	0.0921

Example 31: Positionally Modified Compositions

The table below shows exemplary positionally modified compositions prepared in accordance with the present invention. Target descriptors are: P=PTEN; S=Survivin; E=eIF4E and are indicated following the antisense strand designation.

	SEQ ID NO.	Composition (5' to 3')
	/ISIS NO.	
	52/345838 (as-P)	UCCUGG _m AUCCUU _m CAC _m CAA _m U _m G _m
	53/338939 (s)	CAUUGGUGAAGGAUCCAGGA
25	60/345839 (as-E)	CCUGG _m A _m UCC _m U _m UCACCAA _m U _m G _m
	53/338939 (s)	CAUUGGUGAAGGAUCCAGGA
	56/345853 (as-E)	AUACUC _m A _m GAA _m G _m GUGUCUU _m C _m U _m
	57/338952 (s)	AGAAGACACCUUCUGAGUAU
	24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
30	25/343868 (s)	GGAGAUCAACAUUUUCAAA
	24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
	25/343868 (s)	GGAGAUCAACAUUUUCAAA
	24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
	25/346287 (s)	<u>GGAGAUCAACAUUUUCAAA</u>

24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
25346287 (s)	<u>GGAGAUCAACAUUUUCAAA</u>
24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
25/352511 (s)	GG _m AG _m AU _m CA _m AC _m AU _m UU _m UC _m AA _m A
5 24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
25/352513 (s)	GG _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m C _m A _m A _m A
24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
25/352511 (s)	GG _m AG _m AU _m CA _m AC _m AU _m UU _m UC _m AA _m A
10 24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A
24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A
24/352505 (as-S)	UUUGA _m AAA _m UGU _m UGA _m UCU _m C _m C _m
15 25/352512 (s)	G _m G _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m C _m A _m A _m A _m
56/345853 (as-E)	AUACUC _m A _m GAA _m G _m GUGUCU _m C _m U _m
57/345857 (s)	AG _m A _m A _m G _m A _m C _m A _m C _m C _m U _m U _m C _m U _m G _m A _m G _m U _m A _m U
20 24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
25/352512 (s)	G _m G _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A _m
24/352506 (as-S)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
25/352513 (s)	GG _m A _m G _m A _m U _m C _m A _m A _m C _m A _m U _m U _m U _m U _m C _m A _m A _m A
25 40/335225 (as-P)	<u>UUUGUCUCU_mG_mG_mUCCUUAACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/335226 (as-P)	<u>UUUGUC_mU_mC_mUGGUCCUUAACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/345711 (as-P)	UUUG _i UCUCUG _i GUCCUUAACU _i U
30 39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/345712 (as-P)	UUU _i G _i UCUCUG _i G _i UCCUUA _i C _i UU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/347348 (as-P)	U _i U _i U _i GUC _i UCU _i GGU _i CCU _i UAC _i U _i U _i
39/308746 (s)	AAGUAAGGACCAGAGACAAA

40/348467 (as-P)	<u>U_lU_lU_lGUC_lUCU_lGGU_lCCU_lUAC_lU_lU_l</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
24/355715 (as-S)	UUUG _l AAAAU _l GUUGAUCUC _l C
25/343868 (s)	GGAGAUCAACAUUUUCAAA
5 40/331426 (as-P)	<u>UUUGUCUCU_lG_lG_lUCCUUACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/331695 (as-P)	<u>UUUGUCUCUGGUCCUUAC_lU_lU_l</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/332231 (as-P)	<u>UUUGUCUCUGGUCCUUACU_lU</u>
10 39/308746 (s)	AAGUAAGGACCAGAGACAAA
24/355712 (as-S)	UUUGA _l AAA _l UGU _l UGA _l UCU _m C _m C _m
25/343868 (s)	GGAGAUCAACAUUUUCAAA
24/353538 (as-S)	UUU _l GAAAAU _l GUU _l GAUCU _l C _l C _s
25/343868 (s)	GGAGAUCAACAUUUUCAAA
15 40/336671 (as-P)	UUUGUCUCUGGUCCUUAC _l U _l U _s
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/336674 (as-P)	UUUGUCUCUGGUCCUU _l AC _l U _l U _s
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/336675 (as-P)	UUUGUCUCUGGUCCUUACUU _s
20 39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/336672 (as-P)	UUUGUCUCUGGUC _l C _l U _l UACUU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/336673 (as-P)	UUUGUCUCUGGU _l C _l C _l UUACUU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
25 40/336676 (as-P)	UUUGUCU _l C _l U _l GGUCCUUACUU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/336678 (as-P)	U _l U _l U _l GUCUCUGGUCCUUACUU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
24/352515 (as-S)	UUUGAAAAUGUUGAU _m C _m U _m C _m C _m
30 25/343868 (s)	GGAGAUCAACAUUUUCAAA
61/330919 (as-P)	<u>UUT_eG_eT_eCUCUGGUCCUUACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
62/330997 (as-P)	<u>T_eT_eT_eGTCUCUGGUCCUUACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA

40/333749 (as-P)	<u>UUUGUCUCUGGUCCU_fU_fA_fCUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/333750 (as-P)	<u>UUUGUCUCUGGU_fC_fC_fUUACUU</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
5 40/333752 (as-P)	UUUGUC _f U _f C _f UGGUCCUUACUU
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/333756 (as-P)	<u>UUUGUCUCUGGUCCUUAC_fU_fU_f</u>
39/308746 (s)	AAGUAAGGACCAGAGACAAA
40/334253 (as-P)	<u>UUUGUCUCU_fG_fG_fUCCUUACUU</u>
10 39/308746 (s)	AAGUAAGGACCAGAGACAAA
24/353539 (as-S)	U _t U _t U _t GAAAAU _t GUU _t GAUCU _m C _m C _m
25/343868 (s)	GGAGAUCAACAUUUUCAAA

The above constructs were tested in HeLa cells, MH-S cells or U-87 MG cells using methods described herein. Resulting IC₅₀'s were calculated and are shown below. Also shown are the species to which the compounds were targeted and the cell line in which they were assayed.

	Construct	Species/Cell Line	Gene	IC50
	345838:338939	Mouse/MH-S	eIF4E	0.022859
	345839:338939	Mouse/MH-S	eIF4E	0.01205
20	345853:338952	Mouse/MH-S	eIF4E	0.075517
	352505:343868	Human/HeLA	Survivin	0.17024
	352506:343868	Human/HeLA	Survivin	0.055386
	352506:346287	Human/HeLA	Survivin	0.11222
	352505:346287	Human/HeLA	Survivin	0.96445
25	352505:352511	Human/HeLA	Survivin	0.21527
	352505:352513	Human/HeLA	Survivin	0.12453
	352506:352511	Human/HeLA	Survivin	0.045167
	352505:352514	Human/HeLA	Survivin	0.47593
	352506:352514	Human/HeLA	Survivin	0.11759
30	352506:352514	Human/HeLA	Survivin	0.376
	352506:352514	Human/U-87 MG	Survivin	0.261
	352505:352512	Human/HeLA	Survivin	0.075608
	345853:345857	Mouse/MH-S	eIF4E	0.025677
	352506:352512	Human/HeLA	Survivin	0.11093

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	352506:352513	Human/HeLA	Survivin	0.24503
	335225:308746	Human/HeLA	PTEN	0.809
	335226:308746	Human/HeLA	PTEN	1.57
	308746:345711	Human/HeLA	PTEN	1.13
5	308746:345712	Human/HeLA	PTEN	0.371
	308746:347348	Human/HeLA	PTEN	0.769
	308746:348467	Human/HeLA	PTEN	18.4
	355715:343868	Human/HeLA	Survivin	0.020825
	331426:308746	Human/HeLA	PTEN	0.5627
10	331695:308746	Human/HeLA	PTEN	0.27688
	332231:308746	Human/HeLA	PTEN	5.58
	355712:343868	Human/HeLA	Survivin	0.022046
	353538:343868	Human/HeLA	Survivin	0.491
	353538:343868	Human/U87-MG	Survivin	0.46
15	336671:308746	Human/HeLA	PTEN	0.273
	336674:308746	Human/HeLA	PTEN	0.363
	336675:308746	Human/HeLA	PTEN	0.131
	336672:308746	Human/HeLA	PTEN	0.428
	336673:308746	Human/HeLA	PTEN	0.122
20	336676:308746	Human/HeLA	PTEN	7.08
	336678:308746	Human/HeLA	PTEN	0.144
	352515:343868	Human/HeLA	Survivin	0.031541
	330919:308746	Human/HeLA	PTEN	29.4
	330997:308746	Human/HeLA	PTEN	3.39
25	333749:308746	Human/HeLA	PTEN	1.3
	333750:308746	Human/HeLA	PTEN	0.30815
	333752:308746	Human/HeLA	PTEN	1.5416
	333756:308746	Human/HeLA	PTEN	1.0933
	334253:308746	Human/HeLA	PTEN	0.68552
30	353539:343868	Human/HeLA	Survivin	0.13216

Example 32: Suitable positional compositions of the invention

The following table describes some suitable positional compositions of the invention. In the listed constructs, the 5'-terminal nucleoside or the sense (upper) strand is hybridized to the 3'-terminal nucleoside of the antisense (lower) strand.

5

Compound (sense/antisense)	Construct (sense 5'→3' / antisense)
sense RNA 4'thio (bold) dispersed antisense	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-XXX ₁₇ XXXXX ₁₂ XXX ₉ XXXXXX ₃ X ₂ X ₁ -5'
Sense RNA 2'-OMe (italic)/ 4'-thio (bold) dispersed antisense	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-X ₁₉ X ₁₈ X ₁₇ XXXXXXXXXXXXXXXXXXXX-5'
Sense RNA Chimeric 2'-OMe (italic)/2'- fluoro(bold italic) antisense	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-XXXXXXXXXXXXXXXXXXXX-5'
Alternate MOE(underline)/OH sense Chimeric OMe (italic) / OH antisense	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-X ₂₀ X ₁₉ X ₁₈ XXXXXXXXX ₁₁ X ₁₀ XXX ₇ X ₆ XXXXX-5'
OMe Gapmer Sense / Chimeric OMe (italic) / OH antisense	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-X ₂₀ X ₁₉ X ₁₈ XXX ₁₅ XXX ₁₂ XXXXXX ₆ XXXXX-5'
Sense RNA Chimeric OMe/OH antisense.	5'-XXXXXXXXXXXXXXXXXXXX-3' 3'-XXX ₁₇ XXX ₁₄ XXX ₁₁ XXX ₈ XXX ₅ XXXX-5'

Example 33: Alternating 2'-O-Methyl/2'-F 20mer siRNAs Targeting PTEN in T-24 cells

A dose response experiment was performed in the PTEN system to examine the positional effects of alternating 2'-O-Methyl/2'-F siRNAs. The nucleosides are annotated as to
10 chemical modification as per the legend at the beginning of the examples.

SEQ ID NO. **Composition (5' to 3')**

/ISIS NO.

40/303912 (as) UUUGUCUCUGGUCCUUACUU

39/308746 (s) P-AAGUAAGGACCAGAGACAAA

15 40/340569 (as) P-U_fU_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m

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39/340573 (s)	P-A _f A _m G _f U _m A _f A _m G _f G _m A _f C _m C _f A _m G _f A _m G _f A _m C _f A _m A _f A _m
40/340569 (as)	P-U _f U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m
39/340574 (s)	P-A _m A _f G _m U _f A _m A _f G _m G _f A _m C _f C _m A _f G _m A _f G _m A _f C _m A _f A _m A _f
40/340569 (as)	P-U _f U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m
5 39/308746 (s)	P-AAGUAAAGGACCAGAGACAAA
40/340570 (as)	P-U _f U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m
39/340573 (s)	P-A _f A _m G _f U _m A _f A _m G _f G _m A _f C _m C _f A _m G _f A _m G _f A _m C _f A _m A _f A _m
40/340570 (as)	P-U _f U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m
39/340574 (s)	P-A _m A _f G _m U _f A _m A _f G _m G _f A _m C _f C _m A _f G _m A _f G _m A _f C _m A _f A _m A _f
10 40/340570 (as)	P-U _f U _m U _f G _m U _f C _m U _f C _m U _f G _m G _f U _m C _f C _m U _f U _m A _f C _m U _f U _m
39/308746 (s)	P-AAGUAAAGGACCAGAGACAAA

The above siRNA constructs were assayed to determine the effects of the full alternating 2'-O-methyl/2'-F antisense strands (PO or PS) where the 5'-terminus of the antisense strands are 2'-F modified nucleosides with the remaining positions alternating. The sense strands were prepared with the positioning of the modified nucleosides in both orientations such that for each siRNA tested with 2'-O-methyl modified nucleosides beginning at the 3'-terminus of the sense strand another identical siRNA was prepared with 2'-F modified nucleosides beginning at the 3'-terminus of the sense strand. Another way to describe the differences between these two siRNAs is that the register of the sense strand is in both possible orientations with the register of the antisense strand being held constant in one orientation. Activity of the constructs (at 150 nM) is presented below as a percentage of untreated control.

siRNA	Activity (% untreated control 150 nM)		
Construct		Sense	Antisense
308746/303912	28%	PO unmodified RNA	PS unmodified RNA
25 340574/340569	46%	PO (2'-F, 3'-0)	PO (2'-F, 5'-0)
340574/340570	62%	PO (2'-F, 3'-0)	PS (2'-F, 5'-0)
340573/340569	84%	PO (2'-O-methyl, 3'-0)	PO (2'-F, 5'-0)
340573/340570	23%	PO (2'-O-methyl, 3'-0)	PS (2'-F, 5'-0)
308746/340569	23%	PO unmodified RNA	PO (2'-F, 5'-0)
30 308746/340570	38%	PO unmodified RNA	PS (2'-F, 5'-0)

Within the alternating motif for this assay the antisense strands were prepared beginning with a 2'-F group at the 5'-terminal nucleoside. The sense strands were prepared with the alternating motif beginning at the 3'-terminal nucleoside with either the 2'-F modified nucleoside or a 2'-O-methyl modified nucleoside. The siRNA constructs were prepared with the

internucleoside linkages for the sense strand as full phosphodiester and the internucleoside linkages for the antisense strands as either full phosphodiester or phosphorothioate.

Example 34: Effect of modified phosphate moieties on alternating 2'-O-methyl/2'-F siRNAs

5 Targeting eIF4E

A dose response was performed targeting eIF4E in HeLa cells to determine the effects of selected terminal groups on activity. More specifically the reduction of eIF4E mRNA in HeLa cells by 19-basepair siRNA containing alternating 2'-OMe/2'-F modifications is shown in this example. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. 5'-P(S) is a 5'-thiophosphate group (5'-O-P(=S)(OH)OH), 5'-P(H) is a 5'-H-phosphonate group (5'-O-P(=O)(H)OH) and 5'-P(CH₃) is a methylphosphonate group (5'-O-P(=O)(CH₃)OH). All of the constructs in this assay were full phosphodiester linked.

HeLa cells were plated at 4000/well and transfected with siRNA in the presence of LIPOFECTIN™ (6μL/mL OPTI-MEM) and treated for about 4 hours, re-fed, lysed the following day and analyzed using real-time PCR methods as described herein. The maximum % reduction is the amount of mRNA reduction compared to untreated control cells at the highest concentration (100 nM), with IC50 indicating the interpolated concentration at which 50% reduction is achieved.

SEQ ID NO	SEQUENCES 5'-3' targeted to eIF4E
20 /ISIS NO	
26/341391 (as)	UUGUCUCUGGUCCUUACUU
27/341401 (s)	AAGUAAGGACCAGAGACAA
58/342744 (as)	UCUUAUCACCUUUAGCUCU
59/342764 (s)	AGAGCUAAAGGUGAUAAGA
25 58/351831 (as)	U _m C _f U _m U _f A _m U _f C _m A _f C _m C _f U _m U _f U _m A _f G _m C _f U _m C _f U _m
59/351832 (s)	A _f G _m A _f G _m C _f U _m A _f A _m A _f G _m G _f U _m G _f A _m U _f A _m A _f G _m A _f
58/368681 (as)	P-U _m C _f U _m U _f A _m U _f C _m A _f C _m C _f U _m U _f U _m A _f G _m C _f U _m C _f U _m
59/351832 (s)	A _f G _m A _f G _m C _f U _m A _f A _m A _f G _m G _f U _m G _f A _m U _f A _m A _f G _m A _f
58/379225 (as)	P(S)-U _m C _f U _m U _f A _m U _f C _m A _f C _m C _f U _m U _f U _m A _f G _m C _f U _m C _f U _m
30 59/351832 (s)	A _f G _m A _f G _m C _f U _m A _f A _m A _f G _m G _f U _m G _f A _m U _f A _m A _f G _m A _f
58/379712 (as)	P(H)-U _m C _f U _m U _f A _m U _f C _m A _f C _m C _f U _m U _f U _m A _f G _m C _f U _m C _f U _m
59/351832 (s)	A _f G _m A _f G _m C _f U _m A _f A _m A _f G _m G _f U _m G _f A _m U _f A _m A _f G _m A _f
58/379226 (as)	P(CH ₃)-U _m C _f U _m U _f A _m U _f C _m A _f C _m C _f U _m U _f U _m A _f G _m C _f U _m C _f U _m
59/351832 (s)	A _f G _m A _f G _m C _f U _m A _f A _m A _f G _m G _f U _m G _f A _m U _f A _m A _f G _m A _f

Double stranded construct Activity

	Antisense	Sense	% Control (100 nM)	IC50 (nM)
	341401	341391	103	n/a (neg control)
	342764	342744	11.0	1.26
5	351832	351831	3.5	0.66
	351832	368681	3.6	0.14
	351832	379225	2.8	0.20
	351832	379712	8.0	2.01
	351832	379226	18.1	8.24

10

Example 35: Assay of selected siRNAs targeting PTEN

The constructs listed below were assayed for activity by measuring the levels of human PTEN mRNA in HeLa cells against untreated control levels. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples. "P(S)-" indicates a thiophosphate group (-O-P(=S)(OH)OH).

	SEQ ID NO	SEQUENCES 5'-3' targeted to PTEN
	/ISIS NO	
	26/371789 (as)	P-UUGUCUCUGGUCCUUACUU
	27/341401 (s)	P-AAGUAAGGACCAGAGACAA
20	26/383498 (as)	<u>U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m</u>
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/381671 (as)	P- <u>U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m</u>
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/382716 (as)	P(S)- <u>U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m</u>
25	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/381672 (as)	P- <u>U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m</u>
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/384758 (as)	P(S)-U _t U _t GUCU _m C _m UGG _m U _m CCUUAC _m U _m U _m
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
30	26/384759 (as)	P(S)-U _t U _t GUCU _m C _m UGG _m U _m CCUUAC _m U _m U _m
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/384760 (as)	P(S)-U _t U _t GUCUCUGG _m U _m CCUUAC _m U _m U _m
	27/359351 (s)	A _e A _e G _e UAAGGACCAGAGAC _e A _e A _e
	26/384761 (as)	P(S)-U _t U _t GUCUCUGG _m U _m CCUUAC _m U _m U _m

- 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 26/359455 (as) UUGUCUCUGGUCCUUACUU
- 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 26/384754 (as) P(S)-UUGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 5 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 26/384755 (as) P(S)-U_tU_tGUCUCUGGUCCUUAC_mU_mU_m
- 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 26/384756 (as) P(S)-U_tU_tGUCUCUGGUCCUUAC_mU_mU_m
- 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 10 26/384757 (as) U_tU_tGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 27/359351 (s) A_eA_eG_eUAAGGACCAGAGAC_eA_eA_e
- 26/359455 (as) UUGUCUCUGGUCCUUACUU
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/384754 (as) P(S)-UUGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 15 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/384755 (as) P(S)-U_tU_tGUCUCUGGUCCUUAC_mU_mU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/384756 (as) P(S)-U_tU_tGUCUCUGGUCCUUAC_mU_mU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 20 26/384757 (as) U_tU_tGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/383498 (as) U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/381671 (as) P-U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m
- 25 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/382716 (as) P(S)-U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/381672 (as) P-U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 30 26/384758 (as) P(S)-U_tU_tGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/384759 (as) P(S)-U_tU_tGUCU_mC_mUGG_mU_mCCUUAC_mU_mU_m
- 27/384762 (s) A_eA_eG_eUAAGGACCAGAGAC_tA_tA_t
- 26/384760 (as) P(S)-U_tU_tGUCUCUGG_mU_mCCUUAC_mU_mU_m

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27/384762 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_t A_t A_t$
26/384761 (as)	$P(S)-U_t U_t G U C U C U G G_m U_m C C U U A C_m U_m U_m$
27/384762 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_t A_t A_t$
26/384758 (as)	$P(S)-U_t U_t G U C U_m C_m U G G_m U_m C C U U A C_m U_m U_m$
5 27/366023 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384759 (as)	$P(S)-U_t U_t G U C U_m C_m U G G_m U_m C C U U A C_m U_m U_m$
27/366023 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384760 (as)	$P(S)-U_t U_t G U C U C U G G_m U_m C C U U A C_m U_m U_m$
27/366023 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
10 26/384761 (as)	$P(S)-U_t U_t G U C U C U G G_m U_m C C U U A C_m U_m U_m$
27/366023 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384754 (as)	$P(S)-U U G U C U_m C_m U G G_m U_m C C U U A C_m U_m U_m$
27/359351 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384755 (as)	$P(S)-U_t U_t G U C U C U G G U C C U U A C_m U_m U_m$
15 27/359351 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384756 (as)	$P(S)-U_t U_t G U C U C U G G U C C U U A C_m U_m U_m$
27/359351 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
26/384757 (as)	$U_t U_t G U C U_m C_m U G G_m U_m C C U U A C_m U_m U_m$
27/359351 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$
20 26/359345 (as)	$U_t U_t G U C U C U G G U C C U U A C U_t U_t$
27/384762 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_t A_t A_t$
26/381671 (as)	$U_t U_t G U C U C U G G U C C U U A C_m U_m U_m$
27/384762 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_t A_t A_t$
26/352820 (as)	$P-U_m U_f G_m U_f C_m U_f C_m U_f G_m G_f U_m C_f C_m U_f U_m A_f C_m U_f U_m$
25 27/384762 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_t A_t A_t$
26/352820 (as)	$P-U_m U_f G_m U_f C_m U_f C_m U_f G_m G_f U_m C_f C_m U_f U_m A_f C_m U_f U_m$
27/359351 (s)	$A_e A_e G_e U A A G G A C C A G A G A C_e A_e A_e$
26/384754 (as)	$P(S)-U U G U C U_m C_m U G G_m U_m C C U U A C_m U_m U_m$
27/359351 (s)	$A_f A_m G_f U_m A_f A_m G_f G_m A_f C_m C_f A_m G_f A_m G_f A_m C_f A_m A_f$

30 Double stranded construct Activity

Antisense	Sense	IC50 (nM)
341391	341401	0.152
359980	359351	0.042
384758	359351	0.095

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	384759	359351	0.08
	384760	359351	0.133
	384761	359351	0.13
	384754	359351	0.203
5	384757	359351	0.073
	352820	359351	0.214
	359980	384762	0.16
	384754	384762	0.245
	384755	384762	0.484
10	384756	384762	0.577
	384757	384762	0.131
	384758	384762	0.361
	384759	384762	0.332
	384760	384762	0.566
15	384761	384762	0.362
	359345	384762	0.155
	359346	384762	0.355
	352820	384762	0.474

20 Example 36: Alternating 2'-MOE/2'-OH siRNAs Targeting PTEN

The constructs listed below targeting PTEN were duplexed as shown (antisense strand followed by the sense strand of the duplex) and assayed for activity using methods described herein. The nucleosides are annotated as to chemical modification as per the legend at the beginning of the examples.

25	SEQ ID NO	SEQUENCES 5'-3' targeted to PTEN	IC50 (nM)
	/ISIS NO		
	27/355771 (s)	P-AA _e GU _e AA _e GG _e AC _e CA _e GA _e GA _e CA _e A	273
	40/357276 (as)	P-UUUG _e UCUC _e UGGUCCUU _e ACUU	
	27/355771 (s)	P-AA _e GU _e AA _e GG _e AC _e CA _e GA _e GA _e CA _e A	5.5
30	40/357276 (as)	P-UUUG _e UCUCUGG _e UCCUUACU _e U	

Example 37: Chemically modified siRNA targeted to PTEN: in vivo study

Six- to seven-week old Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected with single strand and double strand compositions targeted to PTEN. The nucleosides

are annotated as to chemical modification as per the legend at the beginning of the examples. Each treatment group was comprised of four animals. Animals were dosed via intraperitoneal injection twice per day for 4.5 days, for a total of 9 doses per animal. Saline-injected animals served as negative controls. Animals were sacrificed 6 hours after the last dose was

5 administered, and plasma samples and tissues were harvested. Target reduction in liver was also measured at the conclusion of the study.

SEQ ID NO SEQUENCES 5'-3' targeted to eIF4E

/ISIS NO

63/116847 C_eT_eG_cC_cT_cAGCCTCTGGAT_cT_cT_cG_cA_c single strand
 10 26/341391 (as) UUGUCUCUGGUCCUUACUU
 27/341401 (s) AAGUAAGGACCAGAGACAA
 26/359995 (as) U_mU_fG_mU_fC_mU_fC_mU_fG_mG_fU_mC_fC_mU_fU_mA_fC_mU_fU_m
 27/359996 (s) A_fA_mG_fU_mA_fA_mG_fG_mA_fC_mC_fA_mG_fA_mG_fA_mC_fA_mA_f

Two different doses of each treatment were tested. Treatment with ISIS 116847, was
 15 administered at doses of 12.5 mg/kg twice daily or at 6.25 mg/kg twice daily.

The siRNA constructs described above (unmodified 341391/341401, 359995/359996 both strands modified) were administered at doses of 25 mg/kg twice daily or 6.25 mg/kg twice daily. Each siRNA is composed of an antisense strand and a complementary sense strand as per previous examples, with the antisense strand targeted to mouse PTEN. ISIS 116847 and all of
 20 the siRNAs of this experiment also have perfect complementarity with human PTEN.

PTEN mRNA levels in liver were measured at the end of the study using real-time PCR and RIBOGREEN™ RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) as taught in previous examples above. Results are presented in the table below as the average % inhibition of mRNA expression for each treatment group, normalized to saline-injected control.

25 **Target reduction by modified siRNAs targeted to PTEN in mouse liver**

Treatment	Dose (mg/kg, administered 2x/day)	% Inhibition	
		Ribogreen	GAPDH
ISIS 116847	12.5	92	95
	6.25	92	95
ISIS 341391/341401	25	12	21
	6.25	2	9
ISIS 359995/359996	25	6	13
	6.25	5	13

As shown in the Table above, all oligonucleotides targeted to PTEN caused a reduction in mRNA levels in liver as compared to saline-treated control. The mRNA levels measured for the ISIS 341391/341401 duplex are also suggestive of dose-dependent inhibition.

The effects of treatment with the RNA duplexes on plasma glucose levels were evaluated in the mice treated as described above. Glucose levels were measured using routine clinical analyzer instruments (eg. Ascencia Glucometer Elite XL, Bayer, Tarrytown, NY). Approximate average plasma glucose is presented in the Table below for each treatment group.

Effects of modified siRNAs targeted to PTEN on plasma glucose levels in normal mice

Treatment	Dose (mg/kg, administered 2x/day)	Plasma glucose (mg/dL)
Saline	N/A	186
ISIS 116847	12.5	169
	6.25	166
ISIS 341391/341401	25	159
	6.25	182
ISIS 359996/359995	25	182
	6.25	169

10

To assess the physiological effects resulting from in vivo siRNA targeted to PTEN mRNA, the mice were evaluated at the end of the treatment period for plasma triglycerides, plasma cholesterol, and plasma transaminase levels. Routine clinical analyzer instruments (eg. Olympus Clinical Analyzer, Melville, NY) were used to measure plasma triglycerides, cholesterol, and transaminase levels. Plasma cholesterol levels from animals treated with either dose of ISIS 116847 were increased about 20% over levels measured for saline-treated animals. Conversely, the cholesterol levels measured for animals treated with either the 25 mg/kg or the 6.25 mg/kg doses of the ISIS 341391/341401 duplex were decreased about 12% as compared to saline-treated controls. The ISIS 359996/359995 duplex did not cause significant alterations in cholesterol levels. All of the treatment groups showed decreased plasma triglycerides as compared to saline-treated control, regardless of treatment dose.

Increases in the transaminases ALT and AST can indicate hepatotoxicity. The transaminase levels measured for mice treated with the siRNA duplexes were not elevated to a level indicative of hepatotoxicity with respect to saline treated control. Treatment with 12.5 mg/kg doses of ISIS 116847 caused approximately 7-fold and 3-fold increases in ALT and AST

25

levels, respectively. Treatment with the lower doses (6.25 mg/kg) of ISIS 116847 caused approximately 4-fold and 2-fold increases in ALT and AST levels, respectively.

At the end of the study, liver, white adipose tissue (WAT), spleen, and kidney were harvested from animals treated with the oligomeric compounds and were weighed to assess gross organ alterations. Approximate average tissue weights for each treatment group are presented in the table below.

Effects of chemically modified siRNAs targeted to PTEN on tissue weight in normal mice

Treatment	Dose (mg/kg, administered 2x/day)	Liver	WAT	Spleen	Kidney
		Tissue weight (g)			
Saline	N/A	1.0	0.5	0.1	0.3
ISIS 116847	12.5	1.1	0.4	0.1	0.3
	6.25	1.1	0.4	0.1	0.3
ISIS 341391/341401	25	1.0	0.3	0.1	0.3
	6.25	0.9	0.4	0.1	0.3
ISIS 359996/359995	25	1.1	0.4	0.1	0.3
	6.25	1.0	0.3	0.1	0.4

As shown, treatment with antisense oligonucleotides or siRNA duplexes targeted to PTEN did not substantially alter liver, WAT, spleen, or kidney weights in normal mice as compared to the organ weights of mice treated with saline alone.

Example 38: Chemically modified siRNA targeted to PTEN: in vivo study

Six- to seven-week old Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected with compounds targeted to PTEN. Each treatment group was comprised of four animals. Animals were dosed via intraperitoneal injection twice per day for 4.5 days, for a total of 9 doses per animal. Saline-injected animals served as negative controls. Animals were sacrificed 6 hours after the last dose of oligonucleotide was administered, and plasma samples and tissues were harvested. Target reduction in liver was also measured at the conclusion of the study.

Two doses of each treatment were tested. Treatment with ISIS 116847 (5'-CTGCTAGCCTCTGGATTTGA-3', SEQ ID NO: 63), a 5-10-5 gapmer was administered at doses of 12.5 mg/kg twice daily or at 6.25 mg/kg twice daily. The siRNA compounds described below were administered at doses of 25 mg/kg twice daily or 6.25 mg/kg twice daily. Each

siRNA is composed of an antisense and complement strand as described in previous examples, with the antisense strand targeted to mouse PTEN. ISIS 116847 and all of the siRNAs of this experiment also have perfect complementarity with human PTEN.

An siRNA duplex targeted to PTEN is comprised of antisense strand ISIS 341391 (5'-
 5 UUGUCUCUGGUCCUACUU-3', SEQ ID NO: 26) and the sense strand ISIS 341401 (5'-
 AAGUAAGGACCAGAGACAA-3', SEQ ID NO: 27). Both strands of the ISIS 341391/341401
 duplex are comprised of ribonucleosides with phosphodiester internucleoside linkages.

Another siRNA duplex targeted to human PTEN is comprised of antisense strand ISIS
 342851 (5'-UUUGUCUCUGGUCCUACUU-3', SEQ ID NO: 40) and the sense strand ISIS
 10 308746 (5'-AAGUAAGGACCAGAGACAAA-3', SEQ ID NO: 39). The antisense strand, ISIS
 342851, is comprised of a central RNA region with 4'-thioribose nucleosides at positions 1, 2, 3,
 5, 16, 18, 19, and 20, indicated in **bold**. The sense strand, ISIS 308746, is comprised of
 ribonucleosides, and both strands of the ISIS 342851/308746 duplex have phosphodiester
 internucleoside linkages throughout.

15 PTEN mRNA levels in liver were measured at the end of the study using real-time PCR
 and RIBOGREEN™ RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) as taught
 in previous examples above. PTEN mRNA levels were determined relative to total RNA or
 GAPDH expression, prior to normalization to saline-treated control. Results are presented in the
 following table as the average % inhibition of mRNA expression for each treatment group,
 20 normalized to saline-injected control.

Target reduction by chemically modified siRNAs targeted to PTEN in mouse liver

Treatment	Dose (mg/kg, administered 2x/day)	% Inhibition	
		Ribogreen	GAPDH
ISIS 116847	12.5	92	95
	6.25	92	95
ISIS 342851/308746	25	11	18
	6.25	7	15
ISIS 341391/341401	25	12	21
	6.25	2	9

As shown in the table, the oligonucleotides targeted to PTEN decreased mRNA levels
 relative to saline-treated controls. The mRNA levels measured for the ISIS 341391/341401
 25 duplex are also suggestive of dose-dependent inhibition.

The effects of treatment with the RNA duplexes on plasma glucose levels were evaluated in the mice treated as described above. Glucose levels were measured using routine clinical analyzer instruments (eg. Ascencia Glucometer Elite XL, Bayer, Tarrytown, NY). Approximate average plasma glucose is presented in the following table for each treatment group.

Effects of chemically modified siRNAs targeted to PTEN on plasma glucose levels in normal mice

Treatment	Dose (mg/kg, administered 2x/day)	Plasma glucose (mg/dL)
Saline	N/A	186
ISIS 116847	12.5	169
	6.25	166
ISIS 342851/308746	25	167
	6.25	173
ISIS 341391/341401	25	159
	6.25	182

To assess the physiological effects resulting from in vivo siRNA targeted to PTEN mRNA, the mice were evaluated at the end of the treatment period for plasma triglycerides, plasma cholesterol, and plasma transaminase levels. Routine clinical analyzer instruments (eg. Olympus Clinical Analyzer, Melville, NY) were used to measure plasma triglycerides, cholesterol, and transaminase levels. Plasma cholesterol levels from animals treated with either dose of ISIS 116847 were increased about 20% over levels measured for saline-treated animals. Conversely, the cholesterol levels measured for animals treated with either the 25 mg/kg or the 6.25 mg/kg doses of the ISIS 341391/341401 duplex were decreased about 12% as compared to saline-treated controls. The other treatments did not cause substantial alterations in cholesterol levels. All of the treatment groups showed decreased plasma triglycerides as compared to saline-treated control, regardless of treatment dose.

Increases in the transaminases ALT and AST can indicate hepatotoxicity. The transaminase levels measured for mice treated with the siRNA duplexes were not elevated to a level indicative of hepatotoxicity with respect to saline treated control. Treatment with 12.5 mg/kg doses of ISIS 116847 caused approximately 7-fold and 3-fold increases in ALT and AST levels, respectively. Treatment with the lower doses (6.25 mg/kg) of ISIS 116847 caused approximately 4-fold and 2-fold increases in ALT and AST levels, respectively.

At the end of the study, liver, white adipose tissue (WAT), spleen, and kidney were harvested from animals treated with the oligomeric compounds and were weighed to assess gross organ alterations. Approximate average tissue weights for each treatment group are presented in the following table.

5 **Effects of chemically modified siRNAs targeted to PTEN on tissue weight in normal mice**

Treatment	Dose (mg/kg, administered 2x/day)	Liver	WAT	Spleen	Kidney
		Tissue weight (g)			
Saline	N/A	1.0	0.5	0.1	0.3
ISIS 116847	12.5	1.1	0.4	0.1	0.3
	6.25	1.1	0.4	0.1	0.3
ISIS 342851/308746	25	1.0	0.3	0.1	0.3
	6.25	0.9	0.4	0.1	0.3
ISIS 341391/341401	25	1.0	0.3	0.1	0.3
	6.25	0.9	0.4	0.1	0.3

As shown, treatment with antisense oligonucleotides or siRNA duplexes targeted to PTEN did not substantially alter liver, WAT, spleen, or kidney weights in normal mice as compared to the organ weights of mice treated with saline alone.

10

Example 39: Stability of alternating 2'-O-methyl/2'-fluoro siRNA constructs in mouse plasma

Intact duplex RNA was analyzed from diluted mouse-plasma using an extraction and capillary electrophoresis method similar to those previously described (Leeds et al., Anal. Biochem., 1996, 235, 36-43; Geary, Anal. Biochem., 1999, 274, 241-248. Heparin-treated mouse plasma, from 3-6 month old female Balb/c mice (Charles River Labs) was thawed from -80 °C and diluted to 25% (v/v) with phosphate buffered saline (140 mM NaCl, 3 mM KCl, 2 mM potassium phosphate, 10 mM sodium phosphate). Approximately 10 nmol of pre-annealed siRNA, at a concentration of 100 µM, was added to the 25% plasma and incubated at 37 °C for 0, 15, 30, 45, 60, 120, 180, 240, 360, and 420 minutes. Aliquots were removed at the indicated time, treated with EDTA to a final concentration of 2 mM, and placed on ice at 0 °C until analyzed by capillary gel electrophoresis (Beckman P/ACE MDQ-UV with eCap DNA Capillary tube). The area of the siRNA duplex peak was measured and used to calculate the percent of intact siRNA remaining. Adenosine triphosphate (ATP) was added at a concentration of 2.5 mM to each injection as an internal calibration standard. A zero time point was taken by diluting

siRNA in phosphate buffered saline followed by capillary electrophoresis. Percent intact siRNA was plotted against time, allowing the calculation of a pseudo first-order half-life. Results are shown in the Table below. ISIS 338918 (UCUUAUCACCUUUAGCUCUA, SEQ ID NO: 54) and ISIS 338943 are unmodified RNA strand with phosphodiester linkages throughout. ISIS 351831 is annotated as $U_mC_fU_mU_fA_mU_fC_mA_fC_mC_fU_mU_fU_mA_fG_mC_fU_mC_fU_m$ and ISIS 351832 as $A_fG_mA_fG_mC_fU_mA_fA_mA_fG_mG_fU_mG_fA_mU_fA_mA_fG_mA_f$ in other examples herein.

Stability of alternating 2'-O-methyl/2'-fluoro siRNA constructs in mouse plasma

Construct	SEQ ID NOs	% Intact siRNA								
		Time (minutes)								
		0	15	30	45	60	120	180	240	360
338918_338943	54 and 55	76.98	71.33	49.77	40.85	27.86	22.53	14.86	4.18	0
351831_351832	58 and 59	82.42	81.05	79.56	77.64	75.54	75.55	75.56	75.55	75

The parent (unmodified) construct is approximately 50% degraded after 30 minutes and nearly gone after 4 hours (completely gone at 6 hours). In contrast, the alternating 2'-O-methyl/2'-fluoro construct remains relatively unchanged and 75% remains even after 6 hours.

Example 40: In vivo inhibition of survivin expression in a human glioblastoma xenograft tumor model

The U-87MG human glioblastoma xenograft tumor model (Kiaris et al., 2000, May-Jun; 2(3):242-50) was used to demonstrate the antitumor activity of selected compositions of the present invention. A total of 8 CD1 nu/nu (Charles River) mice were used for each group. For implantation, tumor cells were trypsinized, washed in PBS and resuspended in PBS at 4×10^6 cells/mL in DMEM. Just before implantation, animals were irradiated (450 TBI) and the cells were mixed in Matrigel (1:1). A total of 4×10^6 tumor cells in a 0.2 mL volume were injected subcutaneously (s.c.) in the left rear flank of each mouse. Treatment with the selected double stranded compositions (dissolved in 0.9% NaCl, injection grade), or vehicle (0.9% NaCl) was started 4 days post tumor cell implantation. The compositions were administered intravenously (i.v.) in a 0.2 mL volume eight hours apart on day one and four hours apart on day two. Tissues (tumor, liver, kidney, serum) were collected two hours after the last dose. Tumors from eight animals from each group were homogenized for western evaluation. Survivin levels were determined and compared to saline controls.

SEQ ID No/ISIS No

Sequence 5'-3'

24/343868 (as)

UUUGAAAAUGUUGAUCUCC

25/343867 (s)	GGAGAUCAACAUUUUCAAA
24/355713 (as)	U _m U _f U _m G _f A _m A _f A _m A _f U _m G _f U _m U _f G _m A _f U _m C _f U _m C _f C _m
25/355714 (s)	G _f G _m A _f G _m A _f U _m C _f A _m A _f C _m A _f U _m U _f U _m U _f C _m A _f A _m A _f
24/353537 (as)	U _t U _t U _t GAAAAUGUUGAUCU _t C _t C _t
5 25/343868 (s)	GGAGAUCAACAUUUUCAAA
24/352506 (as)	UUUGAA _m A _m AUG _m U _m UGAUCU _m C _m C _m
25/352514 (s)	GG _e AG _e AU _e CA _e AC _e AU _e UU _e UC _e AA _e A

Double stranded construct Activity
Antisense Sense % Inhibition of Survivin

10 343868	343867	none
355713	355714	60
353537	343868	48
352506	352514	44

The data demonstrate that modified chemistries can be used to stabilize the constructs
15 resulting in activity not seen with the unmodified construct.

Various modifications of the invention, in addition to those described herein, will be
apparent to those skilled in the art from the foregoing description. Such modifications are also
intended to fall within the scope of the appended claims. Each reference (including, but not
limited to, journal articles, U.S. and non-U.S. patents, patent application publications,
20 international patent application publications, gene bank accession numbers, and the like) cited in
the present application is incorporated herein by reference in its entirety.

What is claimed

1. A composition comprising a first oligomeric compound and a second oligomeric compound, wherein:
 - at least a portion of the first oligomeric compound is capable of hybridizing with at
 - 5 least a portion of the second oligomeric compound;
 - at least a portion of the first oligomeric compound is complementary to and capable of hybridizing to a selected nucleic acid target;
 - one of the first and the second oligomeric compounds comprises nucleosides linked by internucleoside linking groups wherein the linked nucleosides comprise an alternating motif;
 - 10 the other of the first and the second oligomeric compounds comprises nucleosides linked by internucleoside linking groups wherein the linked nucleosides comprise a positionally modified motif or a fully modified motif;
 - the composition further comprising one or more optional overhangings, phosphate moieties, conjugate groups or capping groups.
- 15 2. The composition of claim 1 wherein the oligomeric compound comprising an alternating motif has the formula:

$$5'-A(-L-B-L-A)_n(-L-B)_{nn}-3'$$
 wherein:
 - 20 each L is, independently, an internucleoside linking group;
 - each A is a β -D-ribonucleoside or a sugar modified nucleoside;
 - each B is a β -D-ribonucleoside or a sugar modified nucleoside;
 - n is from about 7 to about 11;
 - nn is 0 or 1; and
 - 25 wherein the sugar groups comprising each A nucleoside are identical, the sugar groups comprising each B nucleoside are identical, the sugar groups of the A nucleosides are different than the sugar groups of the B nucleosides and at least one of A and B is a sugar modified nucleoside.
- 30 3. The composition of claim 2 wherein each A or each B is a β -D-ribonucleoside.
4. The composition of claim 2 wherein each A or each B is a 2'-modified nucleoside wherein the 2'-substituent is selected from halogen, allyl, amino, azido, O-allyl, O-C₁-C₁₀ alkyl, -OCF₃, O-(CH₂)₂-O-CH₃, 2'-O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n) or O-CH₂-C(=O)-N(R_m)(R_n),

where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C_1 - C_{10} alkyl.

5. The composition of claim 4 wherein the 2'-substituent is allyl, O-allyl, O- C_1 - C_{10} alkyl,
15 O- $(CH_2)_2$ -O- CH_3 or 2'-O $(CH_2)_2SCH_3$.
6. The composition of claim 5 wherein the 2'-substituent is O- $(CH_2)_2$ -O- CH_3 .
7. The composition of claim 2 wherein each A and each B is modified nucleoside.
10
8. The composition of claim 7 wherein one of each A and each B comprises 2'-O CH_3
modified nucleosides.
9. The composition of claim 8 wherein the other of each A and each B comprises 2'-F
15 modified nucleosides.
10. The composition of claim 2 wherein the second oligomeric compound comprises an
alternating motif and one of each A and each B are β -D-ribonucleosides.
- 20 11. The composition of claim 10 wherein the other of each A and each B comprises 2'-
modified nucleosides.
12. The composition of claim 11 wherein each 2'-substituent of the 2'-modified nucleosides
is allyl, O-allyl, O- C_1 - C_{10} alkyl, O- $(CH_2)_2$ -O- CH_3 or 2'-O $(CH_2)_2SCH_3$.
25
13. The composition of claim 12 wherein each 2'-substituent of the 2'-modified nucleosides
is O- $(CH_2)_2$ -O- CH_3 .
14. The composition of claim 2 wherein each L is independently a phosphodiester or a
30 phosphorothioate internucleoside linking group.
15. The composition of claim 1 wherein one of the first and the second oligomeric
compounds comprises a fully modified motif wherein essentially each nucleoside of the

oligomeric compound is a sugar modified nucleoside and wherein each sugar modification is the same.

16. The composition of claim 15 wherein each sugar modified nucleoside is selected from
5 2'-modified nucleosides, 4'-thio modified nucleosides, 4'-thio-2'-modified nucleosides and nucleosides having bicyclic sugar moieties.

17. The composition of claim 16 wherein each nucleoside of the fully modified oligomeric compound is a 2'-modified nucleoside.

10

18. The composition of claim 17 wherein each nucleoside of the fully modified oligomeric compound is a 2'-OCH₃ or a 2'-F modified nucleoside.

19. The composition of claim 18 wherein each nucleoside of the fully modified oligomeric
15 compound is a 2'-OCH₃ modified nucleoside.

20. The composition of claim 15 wherein one or both of the 3' and 5'-termini is a β -D-ribonucleoside.

20 21. The composition of claim 1 wherein one of the first and second oligomeric compounds comprises a positionally modified motif.

22. The composition of claim 21 wherein the oligomeric compound comprising a positionally modified motif comprises a continuous sequence of linked nucleosides comprising
25 from about 4 to about 8 regions wherein each region is either a sequence of β -D-ribonucleosides or a sequence of sugar modified nucleosides and wherein the regions are alternating wherein each of the β -D-ribonucleoside regions is flanked on each side by a region of sugar modified nucleosides and each region of sugar modified nucleosides is flanked on each side by a β -D-ribonucleoside region with the exception of regions located the 3' and 5'-termini that will only be
30 flanked on one side and wherein the sugar modified nucleosides are selected from 2'-modified nucleosides, 4'-thio modified nucleosides, 4'-thio-2'-modified nucleosides and nucleosides having bicyclic sugar moieties.

23. The composition of claim 22 comprising from 5 to 7 regions.

24. The composition of claim 22 wherein each of the regions of β -D-ribonucleosides comprises from 2 to 8 nucleosides in length.
25. The composition of claim 22 wherein each of the regions of sugar modified nucleosides
5 comprises from 1 to 4 nucleosides in length.
26. The composition of claim 25 wherein each of the regions of sugar modified nucleosides comprises from 2 to 3 nucleosides in length.
- 10 27. The composition of claim 22 wherein the oligomeric compound comprising a positionally modified motif has the formula:
$$(X_1)_j-(Y_1)_i-X_2-Y_2-X_3-Y_3-X_4$$

wherein :
X₁ is a sequence of from 1 to about 3 sugar modified nucleosides;
15 Y₁ is a sequence of from 1 to about 5 β -D-ribonucleosides;
X₂ is a sequence of from 1 to about 3 sugar modified nucleosides;
Y₂ is a sequence of from 2 to about 7 β -D-ribonucleosides;
X₃ is a sequence of from 1 to about 3 sugar modified nucleosides;
Y₃ is a sequence of from 4 to about 6 β -D-ribonucleosides;
20 X₄ is a sequence of from 1 to about 3 sugar modified nucleosides;
i is 0 or 1; and
j is 0 or 1 when i is 1 or 0 when i is 0.
28. The composition of claim 27 wherein:
25 X₄ is a sequence of 3 sugar modified nucleosides;
Y₃ is a sequence of 5 β -D-ribonucleosides;
X₃ is a sequence of 2 sugar modified nucleosides; and
Y₁ is a sequence of 2 β -D-ribonucleosides.
- 30 29. The composition of claim 28 wherein i is 0 and Y₂ is a sequence of 7 β -D-ribonucleosides.
30. The composition of claim 28 wherein i is 1, j is 0, Y₂ is a sequence of 2 β -D-ribonucleosides and Y₁ is a sequence of 5 β -D-ribonucleosides.

31. The composition of claim 28 wherein i is 1, j is 1, Y₂ is a sequence of 2 β-D-ribonucleosides, Y₁ is a sequence of 3 β-D-ribonucleosides and X₁ is a sequence of 2 sugar modified nucleosides.
- 5 32. The composition of claim 27 wherein each of the sugar modified nucleosides is a 2'-modified nucleoside or a 4'-thio modified nucleoside.
33. The composition of claim 21 wherein the first strand comprises the positional motif.
- 10 34. The composition of claim 1 wherein each of the internucleoside linking groups of the first and the second oligomeric compounds are independently selected from phosphodiester or phosphorothioate.
35. The composition of claim 1 wherein each of the first and second oligomeric compounds
15 independently comprises from about 12 to about 30 nucleosides.
36. The composition of claim 1 wherein each of the first and second oligomeric compounds independently comprises from about 17 to about 23 nucleosides.
- 20 37. The composition of claim 1 wherein each of the first and second oligomeric compounds independently comprises from about 19 to about 21 nucleosides.
38. The composition of claim 1 wherein the first and the second oligomeric compounds form a complementary antisense/sense siRNA duplex.
- 25 39. A method of inhibiting gene expression comprising contacting one or more cells, a tissue or an animal with a composition of claim 1.
40. A method of inhibiting protein levels in a tumor in an animal comprising contacting the
30 animal with a composition of claim 1.
41. The method of claim 40 wherein contacting is via intravenous administration.
42. The method of claim 40 wherein the tumor is a glioblastoma.

43. The method of claim 40 wherein the protein is encoded by the survivin gene.

SEQUENCE LISTING

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Allerson, Charles
Griffey, Richard H.
Swayze, Eric E.

<120> Double Strand Compositions Comprising Differentially
Modified Strands For Use In Gene Modulation

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<220>
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<220>
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<210> 8
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22

<210> 10

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<223> PCR primer

<400> 10

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25

<210> 11

<211> 22

<212> DNA

<213> Artificial Sequence

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<223> PCR probe

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<211> 1842

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<213> H. sapiens

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26

<210> 15

<211> 26

<212> DNA

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<400> 15

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<210> 16

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<212> DNA

<213> M. musculus

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<213> Artificial Sequence

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<223> PCR primer

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19

<210> 18

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<220>
<223> PCR primer

<400> 18
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<210> 19
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<220>
<223> PCR probe

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<223> Oligomeric compound

<400> 20
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<210> 21
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<221> misc_feature
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<221> misc_feature
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<210> 25
<211> 19
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<210> 30

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<210> 31

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<210> 32

<211> 19

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uuacuagaca acuggauau

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<210> 33

<211> 19

<212> RNA

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<223> Oligomeric compound

<400> 33

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<210> 34

<211> 19

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<213> Artificial Sequence

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<223> Oligomeric compound

<400> 34
uuaaaaagug aguagucac 19

<210> 35
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<400> 35
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<210> 36
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<221> misc_feature
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<221> misc_feature
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<400> 38
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<210> 39
<211> 20
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<213> Artificial Sequence

<220>
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<210> 40
<211> 20
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uuugucucug guccuacuu 20

<210> 41
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<400> 41
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<400> 42
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<210> 43
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<400> 43
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<400> 45
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17

<210> 46
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16

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<400> 47
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20

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<210> 50
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<210> 51
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<210> 57
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<400> 57
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<210> 58
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<400> 58
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<210> 60
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<220>
<223> Oligomeric compound

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<210> 61
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<221> misc_feature
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<223> bases at these positions are RNA

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<210> 63
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<212> DNA
<213> Artificial Sequence

<220>
<223> Oligomeric compound

<400> 63
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(54) Title: DOUBLE STRAND COMPOSITIONS COMPRISING DIFFERENTIALLY MODIFIED STRANDS FOR USE IN GENE MODULATION

(57) Abstract: The present invention provides double stranded compositions wherein each strand is modified to have a motif defined by positioning of β -D-ribonucleosides and sugar modified nucleosides. More particularly, the present compositions comprise one strand having an alternating motif and another strand having a hemimer motif, a blockmer motif, a fully modified motif or a positionally modified motif. At least one of the strands has complementarity to a nucleic acid target. The compositions are useful for targeting selected nucleic acid molecules and modulating the expression of one or more genes. In preferred embodiments the compositions of the present invention hybridize to a portion of a target RNA resulting in loss of normal function of the target RNA. The present invention also provides methods for modulating gene expression.

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INTERNATIONAL SEARCH REPORT

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A. CLASSIFICATION OF SUBJECT MATTER

IPC(8) : A61K 48/00; C07H 21/04; C12Q 1/68
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B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
U.S. : 514/44; 435/6, 325, 375; 536/23.1, 24.5

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
WEST, Biosis, Medline, Embase, CA, SciSearch

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WO 2004015107 (GIESE et al.) 19 February 2004 (19.02.2004). Throughout.	1-48
A	WO 2003070918A (BEIGELMAN et al.) 28 August 2003 (28.08.2003). Throughout.	1-43

☐ Further documents are listed in the continuation of Box C.

☐ See patent family annex.

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"O" document referring to an oral disclosure, use, exhibition or other means	
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